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(54) METHODES ET MATERIELS AMELIORES DE TRANSFORMATION

(54) IMPROVED METHODS AND MATERIALS FOR TRANSFORMATION

(57)

Disclosed herein are novel methods and materials directed to transforming a host cell and expressing exogenous RNA therein. Specifically disclosed are DNA-launching platforms used to introduce a replicating viral segment attached to an exogenous polynucleotide into a cell, whereby the exogenous polynucleotide is expressed in said cell and confers a detectable trait.

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- (54) METHODES ET MATERIELS AMELIORES DE TRANSFORMATION
- (54) IMPROVED METHODS AND MATERIALS FOR TRANSFORMATION

(57) L'invention concerne de nouvelles méthodes et de nouveaux matériels destinés à la transformation d'une cellule hôte et à l'expression dans celle-ci d'ARN exogène. Spécifiquement, l'invention concerne des platesformes de lancement d'ADN utilisées pour introduire un segment viral réplicant fixé à un polynucléotide exogène dans une cellule, de manière que le polynucléotide exogène soit exprimé dans ladite cellule et confère un trait détectable.

(57) Disclosed herein are novel methods and materials directed to transforming a host cell and expressing exogenous RNA therein. Specifically disclosed are DNA-launching platforms used to introduce a replicating viral segment attached to an exogenous polynucleotide into a cell, whereby the exogenous polynucleotide is expressed in said cell and confers a detectable trait.



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#### (57) Abstract

Disclosed herein are novel methods and materials directed to transforming a host cell and expressing exogenous RNA therein. Specifically disclosed are DNA-launching platforms used to introduce a replicating viral segment attached to an exogenous polynucleotide into a cell, whereby the exogenous polynucleotide is expressed in said cell and confers a detectable trait.

PCT/US99/11250

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#### DESCRIPTION

# IMPROVED METHODS AND MATERIALS FOR TRANSFORMATION

This invention was made with United States government support awarded by the following agency:

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The United States has certain rights in this invention.

# Background of the Invention

RNA viruses have been found to be valuable tools in the phenotypic and genotypic transformation of targeted cells and tissues. See, e.g., U.S. Patent No. 5,500,360, which teaches novel viral RNA expression vectors. It has been shown that the RNA of the genome of an RNA virus can be modified to include an exogenous RNA segment and that the modified RNA can be introduced into a host cell, replicated therein, and thereby express the exogenous RNA segment.

Current methods of inoculating a host cell with modified RNA viruses involve the in vitro transcription of a particular strand followed by the introduction of the resulting RNA transcripts into the host cell. One problem with the current inoculation method is that the RNA rapidly degrades which causes a low efficiency of infection. In addition, the preparation of the in vitro RNA transcripts is expensive and time consuming.

Further, with the advent of transformation and the genetic engineering of plants, much concern has arisen concerning the potential hazard of the dispersal of dangerous traits into the environment. For example, genes increasing the stress tolerance and/or herbicide resistance of an agriculturally important crop could theoretically "leak" to surrounding less desirable and damaging plants, e.g., through pollen, mechanical or insect dispersal. This phenomenon could create a novel species of "super-weed" which could wreak havoc on the agricultural industry. Existing RNA virus-based vectors can spread to non-target plants by mechanical means and/or by insects. Such spread can be prevented by using vectors that can replicate and/or move only in target plants expressing the appropriate trans-acting factors. Accordingly, there remains a need for less expensive and more efficient methods of transformation of target cells and tissues. Moreover, there is a need for a novel method of transformation which alleviates the potential dangers associated with the unwanted spread of engineered traits into the environment.

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WO 99/61597

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#### Brief Summary of the Invention

The subject invention pertains to improved materials and methods for transforming host cells which involve transfecting said cells with a DNA-launching platform. One aspect of the subject invention pertains to a DNA-launching platform which encodes a modified viral RNA molecule downstream of DNA-dependent RNA polymerase (pol) promoter, whereby the DNAlaunching platform is capable of being introduced into a host cell and effectively "launching" said modified viral RNA molecule into the host cell such that it is replicated and expressed therein. The term "modified viral RNA molecule" as used herein refers to a viral RNA which has been changed from its natural state. Examples of changes of viral RNA include, but are not limited to, removal of a part of viral RNA genome, insertion or substitution of an exogenous RNA, etc. The exogenous RNA segment can be located in a region of the viral RNA molecule such that it does not disrupt the RNA replication. Techniques for such manipulations have been well known to those of ordinary skill in the art for many years. Preferably, the modified viral RNA molecule further comprises a ribozyme which is located in the proximity of the 3' end of the modified viral RNA molecule. The viral segment may have the ability to be replicated with or, alternatively, without the presence of trans-acting viral replicating elements.

Another aspect of the subject invention pertains to a method of genotypically or phenotypically modifying a host cell, comprising introducing a DNA-launching platform which encodes a viral RNA molecule and an exogenous RNA segment in a location which does not disrupt the replication of said viral RNA segment or said exogenous RNA segment, whereby the exogenous RNA segment confers a detectable trait in the host cell. The subject invention applies to a wide array of plant cells.

Still a further aspect of the subject invention pertains to cells in which the DNAlaunching platform of the subject invention has been introduced.

Yet another aspect of the subject invention pertains to a plant comprising cells transfected with the DNA-launching platform.

The novel methods and materials of the subject invention provide a greater inoculation efficiency of RNA viruses because use of DNA-launching platforms of the subject invention are more resistant to degradation than RNA inocula, and because each DNA platform produces multiple RNA transcripts over an extended period of time. As the DNA-launching platform provides a genetically stable in planta archive copy of a desired vector construct, the continuing transcription of said DNA platform will repeatedly reinoculate the host cell with the desired construct. This serves to counteract genetic instability problems that have inhibited the expression of some genes from vectors based on plant and animal RNA viruses. Further, the

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inoculation methods of the subject invention provide a much simpler means of producing inocula in bulk for large scale use, which is cheaper and more efficient than inoculating with *in vitro* RNA transcripts.

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# Brief Description of the Drawings

Figure 1 represents the schematic for producing the 1a and 2a proteins in the host cell.

Figure 2 illustrates an example of an Agrobacterium transformation vector containing an expression cassette capable of expressing 1a and/or 2a BMV proteins.

Figure 3 illustrates several *Agrobacterium* vectors that were produced to transform host plant cells (black rectangles indicate T-DNA borders).

Figure 4 represents the general mechanism of BMV RNA3 launching, and replication.

Figure 5 depicts DNA-launching platforms which can be used in accord with the teachings contained herein. The BMV and CCMV designations denote cis-acting elements.

Figure 6 depicts DNA-launching platforms which can be used in accord with the teachings contained herein.

Figure 7 depicts DNA-launching platforms which can be used in accord with the teachings contained herein.

Figure 8 depicts DNA-launching platforms which can be used in accord with the teachings contained herein.

Figure 9 depicts Agrobacterium vector for delivery of DNA-launching platforms to plant cells (open triangles represent T-DNA borders).

Figure 10 depicts DNA-launching platforms which can be used in accord with the teachings contained herein.

# 25 Legend For Figures 5-10:

35S = CaMV35S promoter

t = termination/polyA + sequences

Rz = ribozyme

NOS = NOS promoter

30 OOA = origin of assembly

FG = foreign gene

Figure 11 shows that BMV replication factors support efficient RNA3 replication in protoplasts.

Figure 12 shows the efficient replication of launched BMV RNA3 in protoplasts.

Figure 13 shows transgenic expression of BMV 1a and 2a mRNAs in N. tabacum and N. benthamiana.

Figure 14 shows the efficient replication of launched BMV RNA3 in (la + 2a)-transgenic plants.

Figure 15 shows the successful GUS expression from the launched BMV RNA3 in (1a + 2a)- transgenic plants.

Figure 16 shows the successful GUS expression from the launched BMV RNA3 in protoplasts.

Figure 17 shows the successful GFP expression from the launched BMV RNA3 in (1a + 2a) - transgenic plants.

Figure 18 shows the successful GFP expression from the launched BMV RNA3 in protoplasts.

Figure 19 shows the efficient replication of the launched BMV RNA3 in (1a + 2a)-transgenic N. benthamiana using Agrobacterium inoculation.

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Figure 20 shows the successful GUS expression from the launched BMV RNA3 having the SHMV coat protein in (1a + 2a)-transgenic plants.

Figure 21 shows that launched BMV replicates, moves cell-to-cell, and spreads long distances in (1a+2a)-transgenic plants.

Figure 22 shows transfection of progeny from (1a+2a)-transgenic N. benthamiana with BMV RNA3 DNA-launching platform and localization of the launched RNÁ3 to the roots.

# Brief Description of the Sequences

SEQ ID NO. 1: pB1LR2 – partial nucleotide sequence includes BMV la expression cassette.

SEQ ID NO. 2: pB1LR3 - partial nucleotide sequence includes BMV la expression cassette.

SEQ ID NO. 3: pB2LR4 – partial nucleotide sequence includes BMV 2a expression cassette.

30 SEQ ID NO. 4: pB2LR5 – partial nucleotide sequence includes BMV 2a expression cassette.

SEQ ID NO. 5: pB12LR6 — partial nucleotide sequence includes BMV 1a and 2a expression cassettes.

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SEQ ID NO. 6: pB12LR7 — partial nucleotide sequence includes BMV la and 2a expression cassettes.

SEQ ID NO. 7: pB12LR8 - partial nucleotide sequence includes BMV la and 2a expression cassettes.

SEQ ID NO. 8: pB12LR9 - partial nucleotide sequence includes BMV la and 2a expression cassettes.

#### Detailed Disclosure of the Invention

To facilitate understanding of the invention, certain terms used throughout are herein defined. The term "RNA virus" as used herein means a virus whose genome is RNA in a double-stranded or single-stranded form, the single strand being a (+) strand or (-) strand.

The terms "transfection" or "transfected" as used herein means an introduction of a foreign DNA or RNA into a cell by mechanical inoculation, electroporation, agroinfection, particle bombardment, microinjection, or by other known methods.

The terms "transformation" or "transformed" as used herein means a stable incorporation of a foreign DNA or RNA into the cell which results in a permanent, heritable alteration in the cell. Accordingly, the skilled artisan would understand that transfection of a cell may result in the transformation of that cell.

The term "launched" as used herein refers to a polynucleotide that has been transcribed from a DNA-launching platform, as described herein and, preferably, replicated.

The term "cis-acting element" as used herein denotes that portion of the RNA genome of an RNA virus which must be present in cis, that is, present as a part of each viral strand as a necessary condition for replication of that strand. Virus replication may depend upon the existence of one or more trans (diffusible) elements which interact with the cis-acting element to carry out RNA replication. If trans-acting elements are necessary for replication, they need not be present or coded for on the modified viral RNA provided, but may be made available within the infected cell by some other means. For example, the trans-acting replication functions may be provided by other, unmodified or modified, components of the viral genome transfected into the cells simultaneously with the modified RNA. The same approach can be used for other trans-acting functions including movement protein, coat protein, and other functions. The target cell may also be premodified, for example, cells may have been previously transformed to provide constitutive expression of the trans-acting functions from a chromosome. The cis-acting element is composed of one or more segments of viral RNA which must be present on any RNA molecule that is to be replicated within a host cell by RNA replication. The segment will most

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WO 99/61597 PCT/US99/11250

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likely be the 5' and 3' terminal portions of the viral RNA molecule, and may include other portions and/or virus open reading frames as well. The cis-acting element is accordingly defined in functional terms: any modification which destroys the ability of the RNA to replicate in a cell known to contain the requisite trans-acting elements, is deemed to be a modification in the cisacting element. Conversely, any modification, such as deletion or insertion in a sequence region which is able to tolerate such deletion or insertion without disrupting replication, is a modification outside the cis-acting element. As is demonstrated herein, using the example of BMV which is known and accepted by those skilled in the art to be a functional example from which substantial portions of an RNA virus molecule may be modified, by deletion, insertion, or by a combination of deletion and insertion, without disrupting replication.

"Exogenous RNA" is a term used to describe a segment or component of RNA to be inserted into the virus RNA to be modified, the source of the exogenous RNA segment being different from the RNA virus itself. The source may be another virus, an organism such as a plant, animal, bacteria, virus, or fungus. The exogenous RNA may be a chemically synthesized RNA, derived from a native RNA, or it may be a combination of the foregoing. The exogenous RNA may provide any function which is appropriate and known to be provided by an RNA segment. Such functions include, but are not limited to, a coding function in which the RNA acts as a messenger RNA encoding a sequence which, when translated by the host cell, results in synthesis of a peptide or protein having useful or desired properties; the RNA segment may also be structural, as for example in ribosomal RNA; it may be regulatory, as for example with small nuclear RNAs or anti-sense RNA; or it may be catalytic. One skilléd in the art will understand that the exogenous RNA may encode, for example, a protein which is a key enzyme in a biochemical pathway, which upon expression effects a desirable phenotypic characteristic, such as altering cell metabolism. Further, the exogenous RNA may encode a protein involved in transcriptional regulation, such as zinc finger, winged-helix, and leucine-zipper proteins. A particularly interesting function is provided by anti-sense RNA, sometimes termed (-) strand RNA, which is in fact a sequence complementary to another RNA sequence present in the target cell which can, through complementary base pairing, bind to and inhibit the function of the RNA in the target cell.

The term "non-viral" is used herein in a special sense to include any RNA segment which is not normally contained within the virus whose modification is exploited for replication and expression, and is therefore used synonymously with "exogenous". Accordingly, a gene derived from a different virus species than that which is modified is included within the meaning of the terms "non-viral" and "exogenous" for the purposes of describing the invention. For

example, a non-viral gene as the term is used herein could include a gene derived from a bacterial virus, an animal virus, or a plant virus of a type distinguishable from the virus modified to effect transformation. In addition, a non-viral gene may be a structural gene derived from any prokaryotic or eukaryotic organism.

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In one embodiment, the subject invention concerns a novel method of transfecting a host cell which uses a DNA-launching platform to introduce viral RNA into the cell. The subject invention is directed towards a method of transfection employing a DNA-launching platform which encodes a modified viral RNA molecule comprising an RNA viral component attached to an exogenous RNA component and a DNA-dependent RNA pol promoter. The DNAdependent RNA pol promoter is preferably but not necessarily fused within up to 10 nucleotides of the 5' transcriptional start site of the modified viral RNA molecule, and more preferably within up to 5 nucleotides of the 5' transcriptional start site. Expression of the DNA-launching platform produces transcripts of the modified viral RNA molecule that are then capable of RNA replication in the presence of replication factors, which can be present in the modified viral RNA and/or may be supplied in trans by other means including expression from chromosome or supplied on different launching plasmids. When the modified viral RNA is replicated, the exogenous RNA can be replicated as well. Further, the exogenous RNA can be expressed in the cell, thereby providing a predetermined phenotypic characteristic. In a preferred embodiment, the DNA launching platform further comprises a nucleotide sequence encoding a self-cleavable ribozyme situated proximate to the 3' end of said RNA molecule. As would be readily apparent to those skilled in the art, known ribozymes may be used in accordance with the subject invention. In a preferred embodiment, the ribozyme cleaves the modified RNA viral molecule at the 3' region. The 3' region can consist of up to 30 nucleotides upstream or downstream of the 3' end; and preferably consists of up to 10 nucleotides upstream or downstream of the 3' end. In a more preferred embodiment, the ribozyme cleaves the modified RNA viral molecule precisely at the 3' end. Other known regulatory sequences, e.g., promoters and/or termination sequences, may also be substituted for and/or included on the DNA-launching platform. A suitable restriction site can be introduced proximate to the 3' end of the modified viral RNA molecule sequence and the DNA molecule can be cleaved by an appropriate restriction enzyme prior to transfection. The term "DNA-launching platform" as used herein is intended to mean a DNA molecule, circular or linear, which has a coding region comprising a segment encoding a modified viral RNA segment, and further, which is capable of being delivered into a cell and subsequently transcribed.

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Possible regulatory sequences can include, but are not limited to, any promoter already shown to be constitutive for expression, such as those of viral origin (CaMV 19S and 35S) or so-called "housekeeping" genes (ubiquitin, actin, tubulin) with their corresponding termination/polyA + sequences. Also, seed-and/or developmentally-specific promoters, such as those from plant fatty acid/lipid biosynthesis genes (ACPs, acyltransferases, desaturases, lipid transfer protein genes) or from storage protein genes (zein, napin, cruciferin, conglycinin, phaseolin, or lectin genes, for example), with their corresponding termination/polyA + sequences can be used for targeted expression. In addition, the gene can be placed under the regulation of inducible promoters and their termination sequences so that gene expression is induced by light (rbcS-3A, cab-1), heat (hsp gene promoters) or wounding (mannopine, HGPGs). It is clear to one skilled in the art that a promoter may be used either in native or truncated form, and may be paired with its own or a heterologous termination/polyA + sequence.

In a particularly preferred embodiment, the subject invention is directed toward a method of genotypically or phenotypically modifying a cell comprising the following steps: a) forming a cDNA molecule of a virus RNA, or of at least one RNA component if the RNA virus is multipartite, the viral RNA having been modified to contain a DNA segment encoding a non-viral RNA component situated in a region able to tolerate such insertion without disrupting replication of the RNA product encoded thereby; b) cloning modified cDNA into a DNA-launching platform; and c) transfecting a suitable host cell with said DNA-launching platform. In a most preferred embodiment, the method further comprises pretransforming a plant with trans-acting viral replication factors and/or other trans-acting factors. Such trans-acting factors may include viral movement proteins(s), coat protein(s), viral protease(s), and other structural and nonstructural genes. In addition to stable expression of trans-acting factors, trans-acting factors may be introduced on separate expression plasmids or may be expressed from RNA transcripts. In a preferred embodiment such trans-acting factors do not replicate. Suitable host cells may include protoplasts, cells in suspension, or cells in tissues or whole organisms.

In a specific embodiment intended as an example of the broader teachings herein, the RNA viral segment can be derived from brome mosaic virus (BMV), whereby the DNA-launching platform comprises DNA encoding the RNA3 segment of the virus. Brome mosaic virus (BMV) is a member of the  $\alpha$  virus-like super family of positive-strand RNA viruses of animals and plants, and has a genome divided among three RNAs. RNA1 and RNA2 encode the la and 2a proteins, respectively, which are necessary for a genomic RNA replication and subgenomic mRNA synthesis (see, e.g., U.S. Patent No. 5,500,360, which to the extent not inconsistent herewith, is incorporated herein by reference). These proteins contain three

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domains conserved in all other members of the α virus-like super family. In (109kDa) contains a c-proximal helicase-like domain and an n-proximal domain implicated in RNA capping, and 2a (94kDa) contains a central polymerase-like domain. See, e.g., French and Ahlquist, (1988). Ia and 2a interact with each other and with cell factors to form a membrane bound viral RNA replication complex associated with the endoplasmic reticulums of infected cells. BMV RNA3, a 2.1-kb RNA, encodes the 3a protein (32kDa) and coat protein (20kDa), which are involved in the spread of BMV infection in its natural plant hosts but are dispensable for RNA replication. See U.S. Patent No. 5,500,360. The 3a or coat protein gene of the RNA3 viral segment can be replaced with exogenous RNA, whereby it does not interfere with the replication element. Further, the exogenous RNA segment can be inserted downstream of an additional subgenomic promoter. Still further, cells or tissues can be pretransformed to express 1a, 2a, 3a, and coat protein, or any combination thereof, wherein DNA-launching platforms containing a foreign gene(s) with the necessary cis-acting components is transfected, such that the foreign gene is replicated and/or expressed.

In one embodiment, the host cell is pretransformed with BMV1 or BMV2 such that it is transgenically engineered to express 1a and 2a proteins. Preferably, the 5' and 3' ends of BMV1 and BMV2 are removed such that they are incapable of replication, but can express 1a and 2a to form a viral RNA replication complex associated with the endoplasmic reticulum of the host cell. Subsequent transfection of a DNA-launching platform comprising the RNA3 viral replication segment, as well as the exogenous RNA of interest, can produce the expression of said exogenous RNA while also preventing the undesired and dangerous spread of viral RNA spillage into the environment. That is, because a plant must have all 3 segments to form infectious BMV particle(s), problems associated with the environmentally hazardous escape of foreign genes through mechanical or insect dispersal of RNA virus vectors are avoided. One skilled in the art will readily appreciate that in the example of BMV that DNA-launching platforms could be also derived from either RNA1 or RNA2. For example, the sequence encoding the Ia protein could be replaced with an exogenous RNA; replication would require the expression of la (e.g., separate expression plasmid). In a preferred embodiment, the DNAlaunching platform also comprises a ribozyme situated proximate to the 3' end of the modified RNA3, wherein said ribozyme cleaves the RNA3 at the 3' end. As would be readily apparent to the skilled artisan with the teachings contained herein, viral segments from other known viruses, and/or subviral agents, can be used to formulate DNA-launching platforms of the subject invention. One skilled in the art will appreciate that BMV is merely one representative example of the many viruses suitable for practicing the subject invention. It is widely accepted that

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principles on which the subject invention is based are broadly applicable to a myriad of viruses. Examples of other such viruses include, but are not limited to, alfalfa mosaic virus (AMV), barley stripe mosaic virus, cowpea mosaic virus, cucumber mosaic virus, reoviruses, polio virus, sindbis virus, vesicular stomatitis virus, influenza virus, retroviruses, and cowpea chlorotic mottle virus (CCMV) and any other viruses that replicate through RNA intermediates and from which a cDNA copy can be obtained. Specifically, as the other viruses are further characterized, those of skill in the art will readily appreciate the applicability of the teachings herein to other suitable viruses as well.

The skilled artisan would easily appreciate that known methods of introducing foreign DNA into cells can be used in accordance with the teachings of the subject disclosure. Such methods include, but are not limited to, mechanical inoculation, particle bombardment, agroinfection, electroporation, and microinjection, as well as other known methods.

Various aspects of the invention can be modified as needed, depending upon specific characteristics of the virus selected as the transforming and transfecting agent and of the RNA segment to be inserted. For example, the inserted gene need not be a naturally occurring gene, but may be modified, a composite of more than one coding segment, or it may encode more than one protein. The RNA may also be modified by combining insertions and deletions in order to control the total length or other properties of the modified RNA molecule. The inserted non-viral gene may be either prokaryotic or eukaryotic in origin. The inserted gene may contain its own translation start signals, for example, a ribosomal binding site and start (AUG) codon, or it may be inserted in a manner which takes advantage of one or more of these components preexisting in the viral RNA to be modified. Certain structural constraints must be observed to preserve correct translation of the inserted sequence, according to principles well understood in the art. For example, if it is intended that the exogenous coding segment is to be combined with an endogenous coding segment, the coding sequence to be inserted must be inserted in reading frame phase therewith and in the same translational direction.

It will be understood by those ordinarily skilled in the art that there may exist certain genes whose transfer does not result in obvious phenotypic modification of the recipient cell. Such may occur, for example, if the translation product of the non-viral gene is toxic to the host cell, is degraded or processed in a manner which renders it non-functional or possesses structural features which render it impossible for the host cell to translate in sufficient quantities to confer a detectable phenotype on the transformed cells. However, the invention does not depend upon any specific property of an RNA segment or gene being transferred. Therefore, the possible existence of RNA segments or genes which fail to confer a readily observable phenotypic trait

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on recipient cells or plants is irrelevant to the invention, and in any case will be readily recognizable by those of ordinary skill in the art without undue experimentation.

An exogenous RNA segment may be inserted at any convenient insertion site in any of the cDNA sequences corresponding to a viral RNA, or component RNA of a multipartite RNA virus, provided the insertion does not disrupt a sequence essential for replication of the RNA within the host cell. For example, for a virus whose coat protein is not essential for replication, an exogenous RNA segment may be inserted within or substituted for the region which normally codes for coat protein. As desired, regions which contribute to undesirable host cell responses may be deleted or inactivated, provided such changes do not adversely affect the ability of the RNA to be replicated in the host cell. For many single component and multipartite RNA viruses, a reduction in the rate of normal RNA replication is tolerable and will in some instances be preferred, since the amount of RNA produced in a normal infection is more than enough to saturate the ribosomes of the transformed cell.

Plant cells which are inoculated in culture will normally remain transfected as the cells grow and divide since the RNA components expressed from the DNA-launching platform are able to replicate and thus become distributed to descendant cells upon cell division. Plants regenerated from phenotypically modified cells, tissues, or protoplasts remain phenotypically modified. Similarly, plants transfected as seedlings remain transfected during growth. Optimal timing of application of the transfecting components will be governed by the result which is intended and by variations in susceptibility to the transfecting components during various stages of plant growth.

Many plant RNA viruses are seed transmitted from one generation to the next. This property can be exploited to effect genotypic transformation of a plant. That is to say, the modified RNA remains transmissible from one generation to the next, just as seed-borne virus infections are transmitted from one generation to the next.

Following are examples which illustrate procedures for practicing the invention. These examples should not be construed as limiting. All percentages are by weight and all solvent mixture proportions are by volume unless otherwise noted.

# Example 1 - Construction of Agrobacterium Vectors

Binary vectors for expressing the BMV 1a and 2a proteins in plants were constructed. Starting with the pBI101.2 construct (Clontech, Palo Alto, CA), the GUS gene was removed by first cutting the construct with EcoRI and SnaBI. The overhanging restriction fragment ends

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were filled in by treatment with Klenow fragments and dNTPs. The restriction fragment ends were religated forming the pB101.2LR1.

The 2a expression cassette was inserted into pBI101.2 LR1. First the pBI101.2LR1 was cut with Hind III and dephosphorylated. Next, pB2PA17 (Dinant et al., 1993) was cut with Hind III and the 2a insert was purified using a low melting agarose gel. The restriction fragment ends were ligated forming the pB2LR4 and pB2LR5 (Figures 3c and 3d).

The la expression cassette was inserted into pB1101.2LR1 by first cutting pB1101.2LR1 with SnaBI and dephosphorylated. pB1PA17 (Dinant et al., 1993) was cut with Pstl and the extra nucleotides were removed with T4 DNA polymerase. The la insert was purified using a low melting agarose gel. The restriction fragment ends were ligated forming the pB1LR2 and pB1LR3 vectors (Figures 3a and 3b).

The la expression cassette was inserted into pB2LR4 and pB2LR5 by cutting pB2LR4 or pB2LR5 with SnaBl and dephosphorylated. PB1PA17 (Dinant et al., 1993) was cut with Pstl, and the extra nucleotides were removed with T4 DNA polymerase. The la insert was purified using low melting agarose gel and ligated with the cut pB2LR4 or pB2LR5 vectors to form pB12LR6, pB12LR7, pB12LR8, and pB12LR9 vectors (Figures 3e-3h).

# Example 2 - Construction of DNA-launching Platform for wtRNA3 of BMV and for RNA Derivatives Containing Foreign Sequences

Vector pRT101 (Töpfer et al., 1987) was cut with PpuMI and the restriction fragment ends were filled in with Klenow fragment and dNTPs, and cut with BamHI and dephosphorylated. Vector pB3RQ39 (Ishikawa et al., 1997) was cut with SnaBI and BamHI; the B3 fragment was isolated from a low melting agarose gel. This fragment was ligated to the cut pRT101 thereby forming pB3LR10 (Figure 4). The pB3LR15 (Figure 4) that is a pB3LR10 derivative has the ClaI-KpnI fragment replaced with the corresponding fragment from pB3TP8 (Janda et al., 1987).

PCR was performed on pRT101 to amplify an EcoRV and EcoRI fragment. To create a StuI site instead of a PpuMI site, a one nucleotide deletion was performed during the PCR process. The resulting PCR product was cut with EcoRV and EcoRI and inserted into dephosphorylated pRT101 cut with EcoRV and EcoRI to form pRT101LR11. The pRT101LR11 was cut with StuI and BamHI and dephosphorylated. PB3RQ39 was cut with SnaBI and BamHI and a B3 fragment was isolated using a low melting agarose gel. The fragment was then ligated to pRT101LR11 to form pB3LR12 (Figure 4).

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Another DNA-launching platform was constructed with wtRNA3 of BMV having a partially doubled CaMV35S promoter; thereby forming pB3LR14 and pB3LR16 (Figure 4).

A DNA-launching platform wherein the BMV RNA3 coat protein was replaced with GUS was also constructed. The pB3MI22 (Ishikawa et al., 1997) was cut with ClaI and StuI and a B3GUS insert was isolated. The pB3LR10 or pB3LR14 DNA-launching constructs were cut with ClaI and StuI and dephosphorylated. The B3GUS fragment was then ligated to the cut pB3LR10 or pB3LR14 thereby forming the pB3GUSLR17 and pB3GUSLR18 DNA-launching constructs (Figure 5).

A DNA-launching platform having a BMV RNA3 with a GUS gene insertion wherein the GUS is downstream of an additional BMV subgenomic promoter was constructed. The pB3LR15 construct was cut with Aval and the restriction fragment ends were filled in with Klenow fragment and dNTPs. Construct was then cut with Clal and dephosphorylated. The pB3MI22 was cut with Clal and StuI and a B3GUS fragment was isolated. The isolated B3GUS fragment was then ligated to the cut pB3LR15 construct to form a new construct of pB3GUSCPLR19 (Figure 5).

A BMV RNA3 based DNA-launching platform with a CP gene inserted downstream of an additional cowpea chlorotic mottle virus (CCMV) subgenomic promoter was constructed. The pB3GUSLR17 construct was cut with Stul and Kpnl and dephosphorylated. The pBC3AJ14 (Pacha and Ahlquist, 1991) was cut with Ndel, the ends were blunted by known methods in the art, and then cut with Kpnl. A coat protein fragment was then isolated. The coat protein fragment was then ligated to the cut pB3GUSLR17 to form a new construct of pB3GUSCPLR22 (Figure 5).

A DNA-launching platform was constructed having a subgenomic RNA4. The pB4MK2 (M. Kroll, personal communications) was cut with SnaBl and BamHl and a RNA4 fragment was then isolated. The pRT101LR11 construct was cut with Stul and BamHl and dephosphorylated. The fragment and the cut pRT101LR11 construct were then ligated forming pB4LR20 (Figure 5a).

A DNA-launching platform wherein the BMV coat protein was replaced with GFP was constructed. pEGFP (Clontech, CA) was cut with Notl, filled in with Klenow fragment and dNTPs, cut with Sall, and GFP insert was isolated using low-melting agarose gel. The pB3LR15 was cut with Sall and Stul and dephosphorylated. The GFP fragment was then ligated to the cut pB3LR15 thereby forming the pB3GFPLR48 (Figure 6e).

A DNA-launching platform having a BMV RNA3 with a GFP gene insertion wherein the CP is downstream of an additional CCMV subgenomic promoter was constructed. The

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pBC3AJ14 (Pacha and Ahlquist, 1991) was cut with NdeI and EcoRI and the ends were blunted by known methods in the art. The coat protein fragment was then isolated and ligated into dephosphorylated and blunted pEGFP cut with NotI and Stul forming pEGFPCPLR49. pEGFPCPLR49 was cut with KpnI and the EGFPCP fragment was isolated using low-melting agarose gel. PB3GFPLR48 was cut with KpnI and dephosphorylated. The EGFPCP fragment was then ligated to the cut pB3GFPLR48 thereby forming the pB3GFPCPLR50 (Figure 6a).

An RNA transcription vector wherein the GFP gene is expressed as a translational fusion with BMV 3a was constructed. The pB3TP10 (Pacha and Ahlquist, 1991) was cut with BamHI and dephosphorylated. The GFP fragment was amplified from pEGFP (Clontech, CA) using PCR and the following primers:

### 5'GCAGTCGACGGTACCGCGGGCC3'

and

#### 5'CGCGGCCGCGGATCCTGTACAGCTCG3'.

The amplified product was cut with BamHl and purified using low-melting agarose gel. The GFP fragment was ligated to the cut pB3TP10 forming pB3GFPLR47 (Figure 6d). The pB3GFPLR47 was cut with EcoRI and transcribed using T7 RNA polymerase.

An Agrobacterium vector containing BMV RNA3 DNA-launching platform was constructed. The pB1101.2LR1 was cut with Smal and dephosphorylated. The pB3LR15 was cut with PvuII and the B3 fragment was purified using a low-melting agarose gel. The B3 fragment was then ligated to the cut pB1101.2LR1 thereby forming pB3LR42 (Figure 9).

A DNA-launching platform wherein the BMV RNA3 coat protein was replaced with the SHMV (Sunn hemp mosaic virus) coat protein and the GUS gene was inserted downstream of an additional BMV subgenomic promoter was constructed. The pB3RS4 (Sacher et al., 1988) was cut with Aval, blunted with Klenow fragment and dNTPs, and cut with KpnI. The SHMV coat protein fragment was isolated using a low-melting agarose gel. The pB3GUSLR17 was cut with Stul and KpnI and dephosphorylated. The SHMV coat protein fragment was ligated to the cut pB3GUSLR17 thereby forming pB3GUSCPLR24 (Figure 7).

Other permutations of DNA-launching platforms containing one or more foreign genes and the necessary cis-acting replication signals will be readily appreciated in view of the teachings herein. For examples, see Figures 5-10.

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# Example 3 — Transfection of N. tabacum Protoplasts with DNA-launching Platform Media:

NT1 Medium (1 liter) was made with Gibco-BRL (MS salt, catalog #11118-031), 3ml of 6% KH2PO4, and 0.2  $\mu$ g/ml 2,4D (final concentration). The pH was adjusted to 5.5-5.7 using KOH, and the resulting mixture was autoclaved.

NT1 Plating Medium (1 liter) was made with NT1 medium and 72.86 g mannitol, the pH was adjusted to 5.5-5.7, and the resulting mixture was autoclaved.

Wash Solution (1 liter) was made with 72.86 g mannitol, the pH was adjusted to 5.5, and the resulting mixture was autoclaved.

Electroporation Buffer was made with 0.8% NaCl, 0.02% KCl, 0.02% KH2PO4, 0.11% Na2HPO4, and 0.4M mannitol. The pH was adjusted to 6.5, and the resulting mixture was autoclaved.

Enzyme Solution was made with 0.4M mannitol, and 20mM MES. The pH was adjusted to 5.5, and the resulting mixture was autoclaved.

Growth conditions: Cells (Nicotiana tabacum) were grown at room temperature in NT1 media with constant shaking (about 200 rpm).

Preparation of cultures for digestion: About 2-3 ml of one-week old suspension culture was subcultured into 50 ml of fresh NT1 media 3 days before the enzyme digestion. The culture was maintained at 28°C under constant shaking.

Enzyme digestion: The enzyme digestion solution was prepared containing the following: 1% cellulysin (Calbiochem) and 0.3% macerase (Calbiochem) in the enzyme solution. The pH was adjusted to 5.5 and filter sterilized.

The cells were centrifuged at 800 rpm for 5 min. The supernatant was discarded. About 40 ml of wash solution was added, cells were resuspended and were centrifuged at 800 rpm for 5 min. The supernatant was discarded. The cells were then resuspended in three volumes of enzyme digestion solution, and incubated for 60 min. at room temperature.

Washing: The cells were transferred into 50 ml plastic tube and centrifuged at 800 rpm for 5 min. The supernatant was discarded. The cells were resuspended in 40 ml of wash solution and centrifuged at 800 rpm for 5 min. The supernatant was discarded. The cells were resuspended in 40ml of electroporation buffer and centrifuged at 800 rpm for 5 min. The supernatant was discarded. The cells were resuspended in four volumes of electroporation buffer.

Electroporation: One ml of cells containing the RNA or DNA inocula was transferred into electroporation cuvettes and placed on ice for 10 min. The cells were then mixed and

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electroporated at 500 microF, 250V. The cuvettes were placed on ice for 10 min. The cells were transferred into 10 ml of NT1 plating media.

Incubation and collection of samples: The cells were incubated at room temperature in dark. Samples were collected 24-48 hrs post inoculation.

RNA Analysis: RNA extraction, denaturing 1% agarose gel electrophoresis and Northern blot hybridization were performed by known methods, such as that performed in Rasochova and Miller (1996). Each lane was loaded with equal amounts (approx. 5  $\mu$ g) of total RNA as determined by spectrophotometry and confirmed by ethidium bromide staining of ribosomal RNA before Northern blot hybridization. 1 X 106 cpm/ml of radioactive probe in hybridization buffer was used per hybridization experiment. Replication of RNA3 was confirmed by detection of sgRNA4, thus showing that BMV RNA replication factors 1a and 2a expressed from expression plasmid(s) support efficient replication of RNA3 supplied as *in vitro* transcript (Figure 11) as well as launched from DNA-launching platform (Figure 12).

# 15 Example 4 - Production of Transgenic N. tabacum Plants

Once a desired molecule was constructed in E. coli, the molecule was transferred into Agrobacierium tumefaciens by the freeze-thaw method. Vectors pBILR2, pB2LR4, pB12LR6, and pB12LR7 were all individually used. An Agrobacterium strain LBA 4404 containing an appropriate helper Ti plasmid was grown in 5 ml of YEP medium overnight at 28°C. Two ml of the overnight culture were added to 50 ml YEP medium in a 250-ml flask and shaken vigorously (250 rpm) at 28°C until the culture grew to an OD<sub>500</sub> of 0.5 to 1.0. The culture was chilled on ice. The cell suspension was centrifuged at 3000 g for 5 min, at 4°C. The supernatant solution was discarded. The cells were resuspended in 1 ml of ice-cold 20 mM CaCl, solution. 0.1-ml aliquots were dispensed into prechilled eppendorf tubes. About 1  $\mu$ g of plasmid DNA was added to the cells. The cells were frozen in liquid nitrogen. The cells were thawed by incubating the test tube in a 37°C water bath for 5 min. 1 ml of YEP medium was added to the tube and incubated at 28°C for 2-4 h with gentle shaking to allow the bacteria to express the antibiotic resistance genes. The tubes were centrifuged for 30 s and the supernatant solution was discarded. The cells were resuspended in 0.1 ml YEP medium, plated on a YEP agar plate containing selection antibiotic(s), and incubated at 28°C. Transformed colonies appeared in 2-3 days.

In vitro clonal copies of approximately three week old Nicotina tabacum, Wisconsin No. 38, were used as the source of explants. Leaf explants were prepared from the second and third fully expanded leaves of in vitro cultures. The leaf pieces were cut into 1 cm x 1 cm squares and

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placed upon TB1 (plus 2.0 mg/l 6-benzyl-aminopurine, and 0.1 mg/l-naphthalene acetic acid) media for 24 hours at 25 °C with a 16 hour photo period.

Agrobacterium tumefaciens strain LBA 4404 containing the preselected binary vector was used for plant transformation. Explants were placed in ~10 ml of overnight grown Agrobacterium culture for 30 min. Leaf explants were then blotted on filter paper and placed on TB2 (plus 1.0 mg/l 6-benzyl-aminopurine and 0.1 mg/l -naphthalene acetic acid) media for 4 days, abaxial side down. Explants are then rinsed three times in sterile water, blotted on filter paper, and placed on TB2 media for regeneration with 100 mg/l kanamycin and 400 mg/l carbenicillin at 25°C, 16 hour photo period, abaxial side down. Explants were transferred to fresh TB2 media with 100 mg/l kanamycin and 400 mg/l carbenicillin every 10 to 14 days until plantlets developed. Plantlets typically developed at 10-14 days. Plantlets were cut from the callus and placed on MST media containing 100 mg/l kanamycin and 400 mg/l carbenicillin to induce rooting. Rooted plants were transferred to soil.

TBI (I liter) included 4.30 g MS salts, 100 mg myo-inositol, 1.0 ml Nitsch and Nitsch vitamins, 30 g sucrose, 2 mg BAP, 0.10 mg of NAA, and 8g Noble agar. The media was adjusted to a pH 5.7 and autoclaved.

TB2 (1 liter) included 4.30 g MS salts, 100 mg myo-inositol, 1.0 ml Nitsch and Nitsch vitamins, 30 g sucrose, 1.0 mg BAP, 0.10 mg NAA, and 8 g Noble agar. The media was adjusted to pH 5.7 and autoclaved.

MST (1 liter) included 4.30 g MS salts, 1.0 ml Nitsch and Nitsch vitamins, 30 g sucrose, 100 mg myo-inositol, and 8.5 g Difco agar. The media was adjusted to pH 5.7 and autoclaved.

YEP (100 ml) included 1.0g Bacto-peptone, 1.0 g Bacto-yeast extract, and 0.5 g NaCl. The media was autoclaved.

RNA Analysis: Total RNA extraction, denaturing 1% agarose gel electrophoresis and Northern blot hybridization was performed by known methods, such as that performed in Rasochova and Miller (1996). Each lane was loaded with equal amounts (approx. 5  $\mu$ g) of total RNA as determined by spectrophotometry and confirmed by ethidium bromide staining of ribosomal RNA before Northern blot hybridization. 1 X 106 cpm/ml of radioactive probe in hybridization buffer was used per hybridization experiment. Figure 13a shows the successful expression of BMV 1a and 2a mRNA in transgenic *N. tabacum*.

Example 5 — Transfection of Transgenic N. tabacum Plants with DNA-launching Platform

Precipitation of DNA onto Microcarriers for Particle Bombardment: (Kikkert, 1993).

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Sterilization of Microcarriers: 80 mg of gold microcarriers were resuspended in 1 ml of 70% ethanol, soaked for 15 min., and centrifuged at 13,000 x g for 5 min. The supernatant was carefully removed and discarded. Particles were resuspended in 1 ml of sterile distilled, deionized water and centrifuged at 13,000 x g for 5 min. The supernatant was carefully removed and discarded. Water washing of particles was repeated 2 more times. After final rinse, particles were resuspended in 1 ml of sterile 50% glycerol.

Coating Microcarriers with DNA: The following was sequentially and quickly added:  $5\mu$ I DNA ( $1\mu$ g/ $\mu$ I),  $50\mu$ I of 2.5M CaCl<sub>2</sub>, and  $20\mu$ I of 0.1M Spermidine.

The mixture was incubated for 10 min. on a vortex shaker at room temperature. Particles were pelleted by centrifugation at  $13,000 \times g$  for 5 sec. Supernatant was carefully removed and discarded. Particles were resuspended in  $140 \mu l$  of 70% ethanol and centrifuged at  $13,000 \times g$  for 5 sec. Supernatant was removed and discarded. Particles were resuspended in  $140 \mu l$  of 100% ethanol and centrifuged at  $13,000 \times g$  for 5 sec. Supernatant was removed and discard. Particles were resuspended in  $50 \mu l$  of 100% ethanol.

Young leaves from tobacco plants grown in vitro on agar-solidified MS medium containing 30g/liter sucrose, were bombarded with 5-µl aliquots of resuspended DNA-coated particles using a PDS1000He biolistic gun (DuPont) and 1100 psi rupture disks (Bio-Rad).

RNA Analysis: Total RNA extraction, denaturing 1% agarose gel electrophoresis and Northern blot hybridization was performed by known methods, such as that performed in Rasochova and Miller (1996). Each lane was loaded with equal amounts (approx. 5 µg) of total RNA as determined by spectrophotometry and confirmed by ethidium bromide staining of ribosomal RNA before Northern blot hybridization. 1 X 106 cpm/ml of radioactive probe in hybridization buffer was used per hybridization experiment. Figure 14a shows that the launched BMV RNA3 replicates efficiently in transgenic plants expressing BMV replication factors 1a and 2a and that the launched RNA3 is unable to replicate in the absence of BMV 1a and/or 2a.

# Example 6 - Production of Transgenic N. benthamiana Plants

Once a desired molecule was constructed in *E. coli*, the molecule was transferred into *Agrobacterium tumefaciens*. Vectors pB1LR2, pB2LR4, pB12LR6, and pB12LR7 were all individually used. An *Agrobacterium* strain LBA 4404 containing an appropriate helper Ti plasmid was grown in 5 ml of YEP medium overnight at 28°C. Two ml of the overnight culture were added to 50 ml YEP medium in a 250-ml flask and shaken vigorously (250 rpm) at 28°C until the culture grew to an OD<sub>500</sub> of 0.5 to 1.0. The culture was chilled on ice. The cell suspension was centrifuged at 3000 g for 5 min. at 4°C. The supernatant solution was discarded.

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The cells were resuspended in 1 ml of ice-cold 20 mM CaCl<sub>2</sub> solution. 0.1-ml aliquots were dispensed into prechilled eppendorf tubes. About 1  $\mu$ g of plasmid DNA was added to the cells. The cells were frozen in liquid nitrogen. The cells were thawed by incubating the test tube in a 37°C water bath for 5 min. 1 ml of YEP medium was added to the tube and incubated at 28°C for 2-4 h with gentle shaking to allow the bacteria to express the antibiotic resistance genes. The tubes were centrifuged for 30 s and the supernatant solution was discarded. The cells were resuspended in 0.1 ml YEP medium. The cells were plated on a YEP agar plate containing selection antibiotic(s) and incubated at 28°C. Transformed colonies appeared in 2-3 days.

In vitro clonal copies of approximately five-seven weeks old N. benthamiana were used as the source of explants. Leaf explants were prepared from the second and third fully expanded leaves of in vitro cultures. The leaf pieces were cut into 1cm x 1cm squares and placed upon MS104 media in 100 x 15 mm plates for 24 hours at 23°C with a 16 hour photo period.

Agrobacterium tumefaciens strain LBA 4404 containing the preselected binary vector was used. Explants were placed in ~10ml of overnight grown Agrobacterium culture for 30 min. Leaf explants were then blotted on filter paper and placed abaxial side down on MS104 media for 4 days. Explants were then rinsed three times in sterile water, blotted on filter paper, and placed on MS104 media for regeneration with 300 mg/L kanamycin and 400 mg/L carbenicillin. Explants were transferred to fresh MS104 media with 300 mg/L kanamycin and 400 mg/L carbenicillin every 10-14 days until plantlets developed. Plantlets typically developed at 31-50 days. Plantlets were cut from the callus and placed on MST media plus 300 mg/L kanamycin and 400 mg/L carbenicillin to induce rooting. Rooted plants were transferred to soil.

One liter of MS104 included 4.3 g MS salt mixture, 1.0 ml B5 vitamin solution, 30 g sucrose, 1.0 mg BA, 0.1 mg NAA, and 8.0 g Phytagar. The media was adjusted to pH 5.8 and autoclaved.

100 ml of YEP included 1.0 g Bacto-peptone, 1.0 g Bacto-yeast extract, 0.5 g NaCl. The media was autoclaved.

One liter of MST included 4.3 g MS salt mixture, 1.0 ml Nitsch & Nitsch vitamins, 30 g sucrose, 100 mg myo-inositol, and 8.5 g Phytagar. The media was adjusted to pH 5.7 and autoclaved.

RNA Analysis: Total RNA extraction, denaturing 1% agarose gel electrophoresis and Northern blot hybridization was performed by known methods, such as that performed in Rasochova and Miller (1996). Each lane was loaded with equal amounts (approx. 5 µg) of total RNA as determined by spectrophotometry and confirmed by ethidium bromide staining of ribosomal RNA before Northern blot hybridization. 1 X 106 cpm/ml of radioactive probe in

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hybridization buffer was used per hybridization experiment. Figure 13b shows the successful expression of BMV 1a and 2a mRNA in transgenic N. benthaniana.

# Example 7 - Transfection of Transgenic N. benthamiana Plants

Precipitation of DNA onto Microcarriers for Particle Bombardment: (From Kikkert (1993) "The biolistic PDS 1000/He device", Plant Cell Tiss. And Org. Cult. 33:221-226)

Sterilization of Microcarriers: 80 mg of gold microcarriers were resuspended in 1 ml of 70% ethanol, soaked for 15 min., and centrifuged at 13,000 x g for 5 min. The supernatant was carefully removed and discarded. Particles were resuspended in 1 ml of sterile distilled, deionized water and centrifuged at 13,000 x g for 5 min. The supernatant was carefully removed and discarded. Water washing of particles was repeated 2 more times. After final rinse, particles were resuspended in 1 ml of sterile 50% glycerol.

Coating Microcarriers with DNA: To the 50  $\mu$ l of particles the following was sequentially and quickly added:  $5\mu$ l DNA ( $1\mu$ g/ $\mu$ l),  $50\mu$ l of 2.5M CaCl<sub>2</sub>, and  $20\mu$ l of 0.1M Spermidine.

The mixture was incubated for 10 min. on a vortex shaker at room temperature. Particles were pelleted by centrifugation at 13,000 x g for 5 sec. Supernatant was carefully removed and discarded. Particles were resuspended in 140  $\mu$ l of 70% ethanol and centrifuged at 13,000 x g for 5 sec. Supernatant was removed and discarded. Particles were resuspended in 140  $\mu$ l of 100% ethanol and centrifuged at 13,000 x g for 5 sec. Supernatant was removed and discarded. Particles were resuspended in 50 $\mu$ l of 100% ethanol.

Young leaves from N. benthamiana plants grown in vitro on agar-solidified MS medium containing 30g/liter sucrose, were bombarded with 5-µl aliquots of resuspended DNA-coated particles using a PDS1000He biolistic gun (DuPont) and 1100 psi rupture disks (Bio-Rad).

RNA Analysis: Total RNA extraction, denaturing 1% agarose gel electrophoresis and Northern blot hybridization was performed by known methods, such as that performed in Rasochova and Miller (1996). Each lane was loaded with equal amounts (approx. 5  $\mu$ g) of total RNA as determined by spectrophotometry and confirmed by ethidium bromide staining of ribosomal RNA before Northern blot hybridization. 1 X 106 cpm/ml of radioactive probe in hybridization buffer was used per hybridization experiment. The launched BMV and RNA 3 showed efficient replication (Figure 14b) in transgenic N. benthamiana plants expressing BMV replication factors 1a and 2a and was unable to replicate in the absence of BMV 1a and/or 2a.

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# Example 8 - Transfection of Transgenic Plants with GUS Containing DNA-launching Platform

Transgenic N. tabacum and N. benthaniana plants were produced according to the procedures discussed above. The plants were transfected with a DNA-launching platform containing a GUS gene (Figure 5a) by particle bombardment as described in Examples 5 and 7. The plants were incubated for 3-5 days and then assayed for β-glucuronidase (GUS) activity using 1 mg/ml X-Gluc (5-bromo-4-chloro-3-indolyl glucucuronide) as substrate in 0.1M potassium phosphate buffer, pH 7.0, 50 μM potassium ferrocyanide, and 2% Triton® X-100. Following an overnight incubation at 37°C, cells replicating launched RNA3 derivatives and expressing the GUS reporter gene from a subgenomic RNA4 gave rise to blue spots (Figure 15). The launched RNA3 derivative did not replicate and express GUS reporter gene in the absence of BMV RNA replication factors 1a and 2a (e.g., in wt N. benthamiana and in wt N. tabacum).

# Example 9 - Transfection of Transgenic Plants Expressing BMV 1a, 2a, 3a, and CP

A plant is transformed with BMV 1a, 2a, 3a, and CP genes whereby those genes are stably expressed in said plant. This can be done with the procedures outlined above. Any modifications that would be needed would be readily apparent to those skilled in the art in light of the teachings contained herein. A DNA-launching platform encoding an RNA replicon which contains a foreign gene and necessary BMV or CCMV cis-acting replication signals to replicate said replicon is constructed (Figure 10b). Foreign genes to be included in said replicon could include, for example, a *Bacillus thuringiensis* polynucleotide that codes for a *B.t.* protein. Other sequences would include, *e.g.*, sequences that encode herbicide resistance, or any other known sequence that encodes peptides or proteins having desired qualities in plants.

Alternatively, plants can be transformed to express BMV 1a, 2a, 3a, and a TMV coat protein in place of the BMV coat protein. A DNA-launching platform is then made containing one or more foreign genes and the necessary cis-acting replication signals, either BMV or CCMV, and a TMV origin of assembly (Figures 8a, 8b, and 10a). This launching platform provides a distinct advantage as TMV is a rod-shaped virus which has no strict limit on the size of RNA that can be encapsidated. Alternatively, TMV movement protein can be used in place of BMV3a (Figure 7c). Hybrids between tobamo and bromoviruses were shown to be viable (Sacher et al., 1988; De Jong and Ahlquist, 1992).

Other permutations and combinations of genes pretransformed and those included in the DNA-launching platform will readily be appreciated by the skilled artisan in light of the teachings herein. (See, e.g., Figures 8c, 10b, and 10c).

As indicated above, CCMV subgenomic promoter can be substituted for BMV sequences in a desired DNA-launching platform. Because the sequence of CCMV subgenomic promoter differs from the sequence of BMV subgenomic promoter, the probability of recombination that would result in loss of a foreign gene is much lower in a construct having a combination of these two different promoters.

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In the above examples, trans-acting components may include, but are not limited to, replication factors, components responsible for cell to cell movement, or components such as the coat protein which may be required for long distance spread, viral proteases responsible for post translational processing, or other known trans-acting functions.

Example 10 - Transfection of N. tahacum Protoplasts with GUS Containing DNA-Launching Platforms

N. tabacum protoplasts isolated using the above described methods were inoculated by electroporation with DNA-launching platforms for BMV RNA3 derivatives in the presence or absence of 1a and 2a expression plasmids. BMV RNA3 derivatives contained the GUS gene in place of the coat protein ORF (Figure 5a) (these were inoculated with or without coat protein expression plasmid, Figure 5b), or had the BMVCP gene translated from an additional subgenomic RNA driven from BMV or CCMV subgenomic promoter (Figures 5c and 5d), or had the SHMV coat protein translated from an additional BMV subgenomic RNA (Figure 7b). Protoplasts were collected by centrifugation (800 rpm, 5 min.) 24 hours post inoculation. The chemiluminescent GUS assay was performed using GUS-Light<sup>TM</sup> (Tropix, MA) according to manufacturer's instructions. Protein concentrations were determined using the Bio-Rad protein kit (Bio-Rad Laboratories, Hercules, CA). The GUS values, determined by luminometer, were adjusted to the same total protein concentration. Figures 16a and 16b show successful GUS expression in protoplasts in the presence of trans-acting BMV replication factors 1a and 2a.

# Example 11 – Transfection of N. tabacum Protoplasts with GFP Containing DNA-Launching Platform

N. tabacum protoplasts isolated by using the above described methods were transfected by electroporation with expression plasmids for trans-acting BMV replication factors 1a and 2a and with DNA-launching platforms for RNA3 derivatives having the GFP gene in place of BMV coat protein ORF (Figure 6e), the CP gene translated from an additional subgenomic RNA (Figure 6a) or with an RNA transcript having the GFP expressed as a fusion protein with BMV 3a ORF (Figure 6d). Protoplasts were incubated for 24 hrs and examined for GFP expression

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using a fluorescent microscope. Figure 18 shows the successful expression of GFP in protoplasts.

# Example 12 - Transfection of (1a + 2a)-Transgenic Plants with BMV RNA3-Based DNA-Launching Platform Containing GFP

N. benthamiana plants were transfected using a particle hombardment as described above with a DNA-launching platform for BMV RNA3 having the GFP gene in place of BMV coat protein (Figure 6e). The GFP expression was determined 24 hrs post inoculation using a fluorescent microscope. Figure 17 shows the successful expression of GFP in (1a + 2a)-transgenic N. benthamiana.

# Example 13 - Transfection of (1a + 2a)-Transgenic N. benthamiana with BMV RNA3 DNA-Launching Platform Using Agrobacterium

N. benthamiana plants were inoculated with BMV RNA3 DNA-launching platform using Agrobacterium tumefaciens. Once the desired construct (pB3LR42) was obtained in E. coli it was transferred to A. tumefaciens strain LBA4404 using a thaw-freeze method as described above. The Agrobacterium was grown overnight in 28°C under constant shaking. A single lower leaf of N. benthamiana were punctured with a needle multiple times and submerged in Agrobacterium culture. The plants were grown at 23°C with a 16 hr photoperiod. The inoculated leaves were harvested 14 days post-inoculation. The total RNA extraction and northern blot hybridization were performed as described above. Figure 19 shows replication of launched BMV RNA3 in inoculated (1a + 2a)-transgenic N. benthamiana.

# Example 14 - Transfection of (1a + 2a)-Transgenic Plants with BMV RNA3-Based DNA-Launching Platform Containing GUS and SHMV Coat Protein

N. benthamiana plants were transfected using a particle bombardment as described above with a DNA-launching platform for BMV RNA3 wherein the BMV coat protein was replaced with the SHMV coat protein (Sunn-hemp mosaic virus) and the GUS gene was inserted downstream of an additional BMV subgenomic promoter (Figure 7b). The GUS expression was determined by histochemical GUS assay described above. Figure 20 shows the successful expression of GUS in (1a + 2a)-transgenic plants.

### Example 15 - Movement of Launched BMV RNA 3

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F1 progeny plants from self-fertilized (1a+2a)-transgenic N. benthamiana BP14 were inoculated with BMV RNA3 DNA launching platform using Agrobacterium tumefaciens. Seedlings were germinated on Smurf media containing Kanamycin. Plants were grown at 23°C with a 16 hr photoperiod. Once the desired construct (pB3LR42) was obtained in E. coli it was transferred to A. tumefaciens strain LBA4404 using a thaw-freeze method as described above. The Agrobacterium was grown overnight at 28°C under constant shaking. A single lower leaf of N. benthamiana was punctured with a needle multiple times and submerged in Agrobacterium culture. The inoculated, middle, and upper leaves were harvested 14 days post-inoculation. Total RNA extraction and northern blot hybridization were performed as described above. RNA3 replication was detected in all leaves tested (Fig. 21). It shows that BMV RNA3 is able to replicate, move cell-to-cell and spread long distance in (1a+2a)-transgenic plants.

# Example 16 - Transfection of Progeny From (1a+2a)-Transgenic N. benthamiana With BMV RNA3 DNA-Launching Platform

Progeny plants from self-fertilized (1a+2a)-transgenic N. benthamiana (designated BP14) were inoculated with BMV RNA3 DNA-launching platform using Agrobacterium as described in Example 13. Control plants (non-transgenic N. benthamiana) were inoculated with the sap from BMV infected barley using inoculation buffer composed of 50mM NaPO<sub>4</sub>, pH7.0, and 1% celite. Root samples were harvested 6 weeks post inoculation. RNA extraction and northern blot hybridization were performed as described above. Figure 22 shows that BMV RNA3 replicated to very high levels in roots. In some (1a+2a)-transgenic plants (Figure 22, lanes 2, 5, 6, 7, 8, 10) replication of launched RNA3 dramatically exceeded replication of wild-type BMV in non-transgenic N. benthamiana plants (Figure 22, lane 1). This shows that this system can be used for delivery of RNA, proteins, peptides or other compounds to roots and enables testing of such compounds for various activities, for example, activities directed against root parasites. For example, proteins with anti-nematode activities can be inserted into RNA3 DNA-launching platform using the above described strategies and expressed in roots upon RNA3 replication. Such proteins can be engineered to be expressed in the cytoplasm or alternatively secreted into the surrounding soil.

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#### Example 17 - Barley Stripe Mosaic Virus

Barley stripe mosaic virus (BSMV) has a tripartite genome (RNA alpha, beta, and gamma). These genomic RNAs have an m7Gppp cap at the 5' end and a t-RNA like structure at the 3' end (Jackson and Hunter, 1989).

A DNA-launching plasmid for BSMV RNA alpha, RNA beta, and RNA gamma containing BSMV RNA cDNA is constructed by precisely fusing at its 5' end to a DNA-dependent RNA polymerase promoter and to a self-cleaving ribozyme at its 3' end. A polyadenylation signal may be also included. Alternatively, a convenient restriction site may be engineered at the 3' end of viral cDNAs. Foreign genes or sequences may be expressed in several ways. For example, DNA-launching plasmids based on BSMV RNA beta may contain a foreign gene or sequence expressed in place of ORF beta a.

Transgenic plants having one or more trans-acting factors fused to the DNA-dependent RNA polymerase promoter and terminator are obtained. Such trans-acting factors may include parts of the viral RNA replicase (ORFs alpha a and/or gamma a) or other trans-acting factors. The trans-acting factors are stably expressed in the plant cell or their expression may be induced if an inducible promoter is used. Cis-acting sequences necessary for BSMV RNA replication are removed from transgenes. Alternatively, the full-length RNA alpha is expressed from the chromosome. Alternatively, ORF gamma a including the 5' untranslated region and ORF gamma b from a seed transmitted strain, such as ND18, are also expressed (Edwards, 1995).

A DNA-launching plasmid is constructed containing the DNA-dependent RNA polymerase promoter precisely fused to the 5' end of the BSMV RNA beta, cis-acting elements important for BSMV RNA beta life cycle, such as the 5' and 3' ends, the intercistronic region between the beta a and beta b ORFs (Zhou and Jackson, 1996) and a foreign gene or sequence in place of ORF beta a (coat protein) which is dispensable for BSMV replication and movement (Petty and Jackson, 1990). Such DNA-launching plasmids may lack the internal poly(A) region as this region is dispensable for replication and contain a ribozyme or a convenient restriction site at the 3' end of the modified viral RNA. Alternatively, a DNA-launching plasmid is constructed from RNA gamma in which ORFs gamma a and/or gamma b are replaced with foreign genes or sequences which may also include the triple gene block genes (ORFs beta b, beta c, and beta d) or a heterologous movement protein (TMV 30K, RCNMV 35K).

# Example 18 - Tobacco Mosaic Virus

Tobacco mosaic virus (TMV) has a single-stranded positive sense RNA genome. The 5' end has an m7Gppp cap and the 3' end contains a t-RNA like structure.

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A DNA-launching plasmid is constructed based on TMV RNA containing TMV cDNA precisely fused at its 5' end to a DNA-dependent RNA polymerase promoter and at its 3' end to a self-cleaving ribozyme. A polyadenylation signal may be also included. Alternatively, a convenient restriction site may be engineered at the 3' end. Foreign gene may be expressed from an additional subgenomic RNA by including an additional subgenomic RNA promoter on the (-) strand.

Transgenic plants are obtained having one or more trans-acting factors fused to the DNA-dependent RNA polymerase promoter and terminator. Such factors may include the viral replicase (126K/183K), movement protein (30K), or coat protein (17.6K). At least one cisacting sequence necessary for TMV RNA replication is removed from transgenes. The transacting factors are stably expressed in the plant cell or their expression may be induced if an inducible promoter is used.

A DNA-launching plasmid is constructed containing the DNA-dependent RNA polymerase promoter precisely fused to the 5' end of the TMV cDNA, cis-acting elements important for the TMV life cycle, such as the 5' and 3' ends, origin of assembly, etc., at least one foreign gene or sequence in place of the trans-acting factor that is expressed from the chromosome, and a ribozyme or a convenient restriction site at the 3' end. Alternatively, the foreign gene sequence can be expressed from an additional subgenomic RNA promoter and the sequence coding for the trans-acting factor that is expressed from the transgene can be deleted from the DNA-launching plasmid. Preferably, if the viral replicase proteins are expressed in transgenic plants, the DNA-launching plasmid will have a deletion of nucleotides 3420-4902, which appears to be a region that inhibits replication in trans. (Lewandowski et al., 1998).

#### Example 19 – Potato Virus X

Potato virus X (PVX) has a single-stranded positive sense RNA genome. The 5' end has an m7Gppp cap and the 3' end is polyadenylated. A full-length cDNA clone of PVX has been constructed and infectious RNA transcripts obtained (Hemenway et al., 1990).

A DNA-launching plasmid is constructed based on PVX RNA containing PVX cDNA precisely fused at its 5' end to a DNA-dependent RNA polymerase promoter and having a polyadenylation site at its 3' end. A convenient restriction site may also be included at the 3' end. A foreign gene may be expressed from an additional subgenomic RNA.

Transgenic plants are obtained having one or more trans-acting factors fused to the DNA-dependent RNA polymerase promoter and terminator. Such factors may include the viral RNA polymerase gene (ORF1-147K), coat protein (ORF5-21K), or triple gene block (ORF2-

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25K, ORF3-12K, ORF4-8K). The triple gene block genes can be expressed individually. Alternatively, they can be expressed as negative sense transcripts from which plus sense subgenomic RNA for ORFs 2, 3, and 4 can be transcribed by the viral replicase. Such transgene will have a DNA-dependent RNA polymerase promoter fused to sequence of ORFs 2, 3, and 4 in the minus sense orientation and the transcribed sequence will include a subgenomic RNA promoter. At least one cis-acting sequence necessary for PVX RNA replication is removed from transgenes. The trans-acting factors are stably expressed in the plant cell or their expression may be induced if an inducible promoter is used.

A DNA-launching plasmid is constructed containing the DNA-dependent RNA polymerase promoter precisely fused to the 5' end of the PVX genome, cis-acting elements important for PVX life cycle, such as the 5' and 3' ends, origin of assembly, etc., at least one foreign gene or sequence in place of the trans-acting factor that is expressed from the chromosome and a polyadenylation signal. Alternatively, the foreign gene sequence can be expressed from an additional subgenomic RNA promoter and the sequence coding for the transacting factor that is expressed transgenically can be deleted from the DNA-launching plasmid.

Alternatively, a DNA-launching plasmid is constructed having a DNA-dependent RNA polymerase promoter, polyadenylation site, and the PVX cDNA sequence in which the ORF2 (25K) is replaced with a foreign gene or sequence. Alternatively, the ORF2 is deleted and the foreign gene is expressed from an additional subgenomic RNA promoter. Such a DNA-launching plasmid is inoculated to transgenic plants expressing movement protein from heterologous virus, such as tobacco mosaic virus (TMV 30K), tomato mosaic virus (ToMV 30K), or red clover necrotic mosaic virus (RCNMV 35K).

# Example 20 - Flock House Virus

Flock house virus (FHV) has a genome consisting of two single stranded RNAs. RNA1 encodes protein A, involved in RNA replication, and protein B that is translated from sg RNA3 and is dispensable for RNA replication. RNA2 encodes virion capsid precursor protein alpha. FHV is infectious to insect, plant, mammalian, and yeast cells (Selling et al., 1990; Price et al., 1996).

A DNA-launching plasmid is constructed for FHV RNA1 and RNA2 containing FHV RNA cDNA precisely fused at its 5' end to a DNA-dependent RNA polymerase promoter and at its 3' end to a self-cleaving ribozyme. A polyadenylation signal may be also included. Alternatively, a convenient restriction site may be engineered at the 3' end. Foreign genes or sequences may be expressed in several ways. For example, DNA-launching plasmids based on

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FHV RNA1 may contain a foreign gene or sequence expressed from subgenomic RNA3 as ORF B replacement or as a translational fusion with ORF B. Alternatively, a foreign gene may be expressed from an additional sg RNA. DNA-launching plasmids based on FHV RNA2 may contain a foreign gene(s) or sequence(s) expressed as a part of polyprotein alpha. Foreign gene(s) in such construct may include sequences necessary for polyprotein clevage. DNA-launching plasmids will preferably also express a movement protein of a heterologous plant virus, such as 30K of TMV or 35K of RCNMV. Alternatively, DNA-launching plasmids will be inoculated onto transgenic plants expressing such movement protein.

Transgenic plants are obtained having one or more trans-acting factors fused to the DNA-dependent RNA polymerase promoter and terminator. Such factors may include protein A or capsid protein precursor alpha, and preferably will also include a movement protein from a plant virus, such as 30K of TMV or 35K of RCNMV. Trans-acting factors are stably expressed in the plant cell or their expression may be induced if an inducible promoter is used. Transgenically expressed trans-acting factors preferably lack at least one cis-acting factor which is necessary for their replication, such as the 5' and/or 3' end.

A DNA-launching plasmid is constructed based on FHV RNA1 or FHV RNA2 containing a DNA-dependent RNA polymerase promoter precisely fused to the 5' end of RNA1 (or RNA2), cis-acting elements important for FHV RNA1 (or RNA2) replication, such as the 5' and 3' ends, at least one foreign gene or sequence and a self-cleaving ribozyme at the 3' end. Polyadenylation signal may also be included. Alternatively, a convenient restriction site may be engineered at the 3' end of the modified viral RNA sequence of the DNA-launching plasmid. DNA-launching plasmids based on FHV RNA1 may contain a foreign gene or sequence in place of ORF A. Alternatively, the ORF A may be deleted and the foreign gene may be expressed from subgenomic RNA3, for example as an ORF B replacement or as a translational fusion with ORF B. Alternatively, DNA-launching plasmid may contain two exogenous RNA sequences, one in the place of ORF A and the other expressed from the subgenomic RNA3. DNA-launching plasmids based on FHV RNA2 may contain a foreign gene(s) or sequence(s) in place of ORF alpha or expressed as a part of polyprotein alpha. Foreign gene(s) in such a construct may include sequences necessary for polyprotein clevage.

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# Example 21 - Tomato Spotted Wild Virus

Tomato spotted wild virus (TSWV) is a tripartite (RNA L, M, S), negative sense and ambisense, single stranded RNA virus.

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Transgenic plants are obtained having one or more trans-acting factors fused to the DNA-dependent RNA polymerase promoter and terminator. Such factors include the putative TSWV polymerase gene (ORF L), ORF N, and possibly other trans-acting factors (NSm or NSs). At least one cis-acting sequence, such as 5' and/or 3' ends, which are necessary for TSWV RNA replication are removed from the transgene. Trans-acting factors are stably expressed in the plant cell or their expression may be induced if an inducible promoter is used.

A DNA-launching plasmid is constructed based on TSWV RNA M in which the G1 and G2 coding sequences are replaced with at least one foreign gene or sequence. Such DNA-launching plasmid contains a DNA-dependent RNA polymerase promoter and TSWV RNA M cDNA fused to the self-cleaving ribozymes at the 5' and 3' ends. Alternatively, a DNA-launching plasmid is constructed based on TSWV RNA S in which the N coding region is replaced with a foreign gene or sequence.

# Example 22 - Barley Mild Mosaic Virus

Genome of barley mild mosaic virus (BaMMV) consists of two positive sense, single-stranded, 3'-polyadenylated RNAs. The RNA1 encodes proteins related to the potyviral P3, 6K1, CI, 6K2, NIa-VPg, NIa-Pro, NIb and capsid protein (Kashiwazaki et al., 1990). The RNA2 encodes P1 and P2 protein (Kashiwazaki et al., 1991). The P1 protein is related to the potyviral HC-Pro and the P2 protein is important for fungal transmission. An isolate was obtained containing a deletion in the P2 protein (Timpe and Kuhne, 1995) thus indicating that P2 is dispensable for viral RNA replication.

A DNA-launching plasmid is constructed for BaMMV RNA1 and RNA2 containing BaMMV RNA cDNA precisely fused at its 5' end to a DNA-dependent RNA polymerase promoter and a polyadenylation site at its 3' end. Foreign genes or sequences may be expressed in several ways. For example, DNA-launching plasmids based on BaMMV RNA2 may contain a foreign gene or sequence expressed as a part of polyprotein which can be cleaved and a foreign protein can be released.

Transgenic plants are obtained having the BaMMV RNA1 cDNA lacking the 5' and 3' ends fused to the DNA-dependent RNA polymerase promoter and terminator.

A DNA-launching plasmid is constructed based on BaMMV (isolate M) RNA2. Such plasmid contains a DNA-dependent RNA polymerase promoter precisely fused to the 5' end of RNA2, RNA2 cis-acting replication signals located in the 5' and 3' ends, P1 ORF and a foreign gene in place of P2 ORF or expressed as a part of P1/P2 polyprotein which can be cleaved and a foreign protein can be released.

The contents of all references cited throughout are incorporated herein by this reference to the extent they are not inconsistent with the disclosure, teachings, and principles of the subject invention.

It should be understood that the examples and embodiments described herein are for illustrative purposes only and that various modifications or changes in light thereof will be suggested to persons skilled in the art and are to be included within the spirit and purview of this application.

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180	TATCTCCTCT	ACCTCTCTCA	CTGTGGTTT	GGCGACGCGC	ACACATTTCC	AGCGATTGAG	4320
	ACTCCAGTCT	TCAGGTATTO	AGACCCCTAC	GTACTTAGAT	ATGTCTTCAA	ACCATACACA	4380
	GTGACGTAGT	GTCTCCCGGG	GGCAGCGTAA	ATTTGTAGCG	ATGATCTTAT	AGGTCATGAT	4440
185	GTTACATTTC	AGCATTTCGC	GCTCCAACAG	ATAGGTGGTT	CCATCGATGC	AATGCACCGA	4500
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190	TTCAAAATCG	AATTTGATCA	CCTCATCCGC	GCCTGACCCG	TCACGTTGCC	AGTGACATTT	4620
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	ACGTAGTACG	CGTACTCCAT	GCGAATGCAT	GGCGTCACAC	AGACCTTGGA	AGCCCATATC	4740
195	ATAACCGCCG	TGGATACAGA	TAGCCCAATC	AGCTTGGACA	TCACAATCTT	GAGCTCGGTT	4800
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200	GCACATCCTC	TCCTCATGTC	GGGCAGCGTC	TCTAACACCC	AACACAGGAC	AACAACTGTG	4920
200	CACCCTTTTA	TCCCTTCTTG	AAAAGTGATG	CCACCAAGAC	CCTCCGAAAT	CTATAACGGG	4980
	GTCTTCAGGG	GGAAAACTGT	CGAGACAGTC	ATAATGCTCC	GCTACACGCA	GAGCACCAGC	5040
205	CAGGCTATGG	GGCGCATGAT	ACTGCTGAGT	CAAATTTAAG	TCAAAGGCAC	CACCATAACG	5100
	GTCACGGAAG	GCGTCAGCCT	CCTCAATAGA	GAGCTTATTG	CGAACGTTGA	TTTTCTTAGA	5160
210	CCTTTTCGCG	TATTCAATCT	GCGCAGATAA	CTGTTGCGCA	ACCTGATTGT	CTACGATGTC	5220
2.0	TTGGGCACTC	TGGCTGTCAG	CACCCTTCTC	AGCAATCAAC	TTCAGCAAAT	CGATAGAACT	5280
	TGACATTTTG	TTGGTGAAAA	ACAAAGAACA	AGTAGCAGAA	CCGTGGTCGA	GGTCCTCTCC	5340
215	AAATGAAATG	AACTTCCTTA	TATAGAGGAA	GGGTCTTGCG	AAGGATAGTG	GGATTGTGCG	5400
	TCATCCCTTA	CGTCAGTGGA	GATATCACAT	CAATCCACTT	GCTTTGAAGA	CGTGGTTGGA	5460
220	ACGTCTTCTT	TTTCCACGAT	GTTCCTCGTG	GGTGGGGGTC	CATCTTTGGG	ACCACTGTCG	5520
220	GTAGAGGCAT	TCTTGAACGA	TAGCCTTTCC	TTTATCGCAA	TGATGGCATT	TGTAGAAGCC	5580

WO 99/61597

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PCT/US99/11250

ATCTTCCTTT TCTACTGTCC TTTCGATGAA GTGACAGATA GCTGGGCAAT GGAATCCGAG 5640 225 GAGGTTTCCC GATATTACCC TTTGTTGAAA AGTCTCAATA GCCCTCTGGT CTTCTGAGAC 5700 TGTATCTTTG ATATTCTTGG AGTAGACGAG AGTGTCGTGC TCCACCATGT TGACCGGGTG 5760 GTCAGTCCCT TATGTTACGT CCTGTAGAAA CCCCAACCCG TGAAATCAAA AAACTCGACG 5820 230 GCCTGTGGGC ATTCAGTCTG GATCGCGAAA ACTGTGGAAT TGATCAGCGT TGGTGGGAAA 5880 GCGCGTTACA AGNAAGCCGG GCAATTGCTG TGCCAGGCAG TTTTAACGAT CAGTTCGCCG 5940 235 ATGCAGATAT TCGTAATTAT GCGGGCAACG TCTGGTATCA GCGCGAAGTC TTTATACCGA 6000 AAGGTTGGGC AGGCCAGCGT ATCGTGCTGC GTTTCGATGC GGTCACTCAT TACGGCAAAG 6060 TGTGGGTCAA TAATCAGGAA GTGATGGAGC ATCAGGGCGG CTATACGCCA TTTGAAGCCG 6120 240 ATGTCACGCC GTATGTTATT GCCGGGAAAA GTGTACAATT CACTGGCCGT CGTTTTACAA 6180 CGTCGTGACT GGGAAAACCC TGGCGTTACC CAACTTAATC GCCTTGCAGC ACATCCCCCT 6240 245 TTCGCCAGCT GGCGTAATAG CGAAGAGGCC CGCACCGATC GCCCTTCCCA ACAGTTGCGC 6300 AGCCTGAATG GCGAATGNNN NNNNAATTCA GTACATTAAA AACGTCCGCA ATGTGTTATT 6360 AAGTTGTCTA AGCGTCAATT TGTTTACACC ACAATATATC CTGCCACCAG CCAGCCAACA 6420 250 GCTCCCCGAC CGGCAGCTCG GCACAAAATC ACCACTCGAT ACAGGCAGCC CATCAGNNNN 6480 255 иминимими имимимими имимимими имимимими имимимими имимимимими 6720 260 имимимим имимимими имимимими имимимими имимимими имимимими 6840 265 имининии имининини имининини имининини имининини имининини 6900 имилимим имимимими имимимими имимимими имимимими имимимими 6960 270 7074

<210> 2

<sup>275 &</sup>lt;211> 6750

<sup>&</sup>lt;212> DNA

<sup>&</sup>lt;213> Brome mosaic virus

	540U2 Z						
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	AGGGAGTCAC	GTTATGACCC	CCGCCGATG	A CGCGGGACA	A GCCGTTTTAC	GTTTGGAACT	120
	GACAGAACCO	G CAACGATTGA	AGGAGCCAC	T CAGCCGCGG	G TTTCTGGAGI	TTAATGAGCT	. 180
285	AAGCACATAC	GTCAGAAACC	ATTATTGCG	C GTTCAAAAG	r cgcctaaggi	CACTATCAGO	240
	TAGCAAATAT	TTCTTGTCAA	AAATGCTCC	A CTGACGTTC	C ATAAATTCCC	CTCGGTATCC	300
290	AATTAGNNNN	<i>א</i> ממממממממ	иииииииии	N GATCGTTTC	G CATGATTGAA	CAAGATGGAT	360
	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTG	G AGAGGCTAT	r cggctatgac	TGGGCACAAC	420
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295	TTTTTGTCAA	GACCGACCTG	TCCGGTGCC	TGAATGAACT	r gcaggacgag	GCAGCGCGGC	540
	TATCGTGGCT	GGCCACGACG	GGCGTTCCT	r GCGCAGCTG1	r GCTCGACGTT	GTCACTGAAG	600
300	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGC	GGATCTCCTG	TCATCTCACC	660
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	ATCCGGCTAC	CTGCCCATTC	GACCACCAAC	G CGAAACATCG	G CATCGAGCGA	GCACGTACTC	780
305	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	A AGAGCATCAG	GGGCTCGCGC	840
	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	CGGCGATGAT	CTCGTCGTGA	900
310	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTCA	960
	TCGACTGTGG	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	CATAGCGTTG	GCTACCCGTG	1020
	ATATTGCTGA	AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG	1080
315	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	TGACGAGTTC	TTCTGANNNN	1140
	имимимими	имимимими	GATCGTTCAA	ACATTTGGCA	ATAAAGTTTC	TTAAGATTGA	1200
320	ATCCTGTTGC	CGGTCTTGCG .	ATGATTATCA	TATAATTTCT	GTTGANTTAC	GTTAAGCATG	1260
	AATTAATTAA	CATGTAATGC .	ATGACGTTAT	TTATGAGAŤG	GGTTTTTATG	ATTAGAGTCC	1320
	CGCAATTATA	CATTTAATAC	GÇGATAGAAA	ACAAAATATA	GCGCGCAAAC '	raggataaat :	1380
325	TATCGCGCGC	GGTGTCATCT	ATGTTACTAG	ATCGGGCCTC	CTGTCAATGC '	reeceeceec :	1440
	TCTGGTGGTG	GTTCTGGTGG (	CGGCTCTGAG	GGTGGTGGCT	CTGAGGGTGG (	CGGTTCTGAG :	1500
330	GGTGGCGGCT	CTGAGGGAGG	CGGTTCCGGT	GGTGGCTCTG	GTTCCGGTGA	TTTTGATTAT :	1560
J = 4	GAAAAGATGG	CAAACGCTAA 1	FAAGGGGGCT	AAGDCDADTA	ATGCCGATGA	AAACGCGCTA .	1620

	CAGTCTGACG	CTAAAGGCAA	ACTTGATTCT	GTCGCTACTG	ATTACGGTGC	TGCTATCGAT	1680
335	GGTTTCATTG	GTGACGTTTC	CGGCCTTGCT	AATGGTAATG	GTGCTACTGG	TGATTTTGCT	1740
	GGCTCTAATT	CCCAAATGGC	TCAAGTCGGT	GACGGTGATA	ATTCACCTTT	AATGAATAAT	1800
340	TTCCGTCAAT	ATTTACCTTC	CCTCCCTCAA	TCGGTTGAAT	GTCGCCCTTT	TGTCTTTGGC	1860
340	CCAATACGCA	AACCGCCTCT	CCCCGCGCGT	TGGCCGATTC	ATTAATGCAG	CTGGCACGAC	1920
	AGGTTTCCCG	ACTGGAAAGC	GGGCAGTGAG	CGCAACGCAA	TTAATGTGAG	TTAGCTCACT	1980
345	CATTAGGCAC	CCCAGGCTTT	ACACTTTATG	CTTCCGGCTC	GTATGTTGTG	TGGAATTGTG	2040
	AGCGGATAAC	AATTTCACAC	AGGAAACAGC	TATGACCATG	ATTACGCCAA	GCTTGCATGC	2100
350	CTGCAGGTCG	ACTCTAGAGG	ATCCCCGGTC	AACATGGTGG	AGCACGACAC	TCTCGTCTAC	2160
330	TCCAAGAATA	TCAAAGATAC	AGTCTCAGAA	GACCAGAGGG	CTATTGAGAC	TTTTCAACAA	2220
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	ATCGTTCAAG	AATGCCTCTA	CCGACAGTGG	TCCCAAAGAT	GGACCCCCAC	CCACGAGGAA	2400
360	CATCGTGGAA	AAAGAAGACG	TTCCAACCAC	GTCTTCAAAG	CAAGTGGATT	GATGTGATAT	2460
300	CTCCACTGAC	GTAAGGGATG	ACGCACAATC	CCACTATCCT	TCGCAAGACC	CTTCCTCTAT	2520
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365	TTTTTCACCA	ACAAAATGTC	AAGTTCTATC	GATTTGCTGA	AGTTGATTGC	TGAGAAGGGT	2640
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370	CAGATTGAAT	ACGCGAAAAG	GTCTAAGAAA	ATCAACGTTC	GCAATAAGCT	CTCTATTGAG	2760
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	ATCTGTATCC	ACGCCGTTA	TGATATGGGC	TTCCAAGGTC	TGTGTGACGC	CATGCATTCG	3180
385	CATGGAGTAC	GCGTACTACG	TGGTACCGTT	ATGTTCGACG	GCGCCATGTT	GTTTGACCGC	3240

WO 99/61597

PCT/US99/11250

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390	GTGATCAAAT	TCGATTTTGA	AAATGAAAGC	ACATTATCTT	ACATCCACGG	ATGGCAAGAT	3360
2.0	TTGGGCTCAT	TTTTCACCGA	GTCGGTGCAT	TGCATCGATG	GAACCACCTA	TCTGTTGGAG	3420
	CGCGAAATGC	TGAAATGTAA	CATCATGACC	TATAAGATCA	TCGCTACAAA	TTTACGCTGC	3480
395	CCCCGGGAGA	CACTACGTCA	CTGTGTATGG	TTTGAAGACA	TATCTAAGTA	CGTAGGGGTC	3540
	TCAATACCTG	AAGACTGGAG	TCTCAATCGC	TGGAAATGTG	TGCGCGTCGC	CAAAACCACA	3600
400	GTGAGAGAGG	TAGAGGAGAT	AGCTTTCAGA	TGTTTCAAGG	AAAGTAAAGA	ATGGACTGAG	3660
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405	TTGACTTTGA	ATCTGTATCA	AAAGTACGAA	AAGCTTACGG	CCCTCCGCGA	TGGGATGGAA	3840
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	TATGCGGACA	GTTTTAAGTT	TCTGACTCGT	CTCTCAAACG	TTGAAGAATT	TGAGCAAGAT	4020
•	TCTGTACCGA	TATCACGTTT	GAGAACGTTT	TGĢACTGAAG	AGGACTTATT	CGACCGGCTG	4080
415	GAGCATGAAG	TGCAGACAGC	CAAGACCAAG	CGCTCGAAGA	AGAAGGCGAA	AGTCCCGCCA	4140
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	TGTAAGAGAT	TACATAACAA	CTCCGAGTCT	AATCTTCGTC	ACCTATGGGA	CATTTCCGGC	4380
425					CCTACCATCG		
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430	GTTGGATATA	ATGAGCATGG	TTTAGGTCCG	AAGCACGCAG	ATGAAACGTA	CATTGTTGAT	4560
	AAAACATGTG	CATGCTCTAA	CTTGAGGGAC	ATTGCAGAAG	CTAGCGCCAA	AGTTTCTGTC	4620
	CCTACATGCG	ATATTTCCAT	GGTTGATGGA	GTTGCGGGAT	GCGGTAAAAC	CACTGCCATA	4680
435	AAAGATGCAT	TCCGTATGGG	AGAGGACCTA	ATTGTGACGG	CGAATCGTAA	ATCGGCCGAG	4740
	GACGTCAGGA	TGGCTTTATT	CCCTGACACT	TATAATTCCA	AGGTAGCTTT	GGACGTTGTG	4800
440	CGCACCGCGG	ATTCTGCGAT	CATGCACGGT	GTACCGTCCT	GTCATAGGCT	GCTTGTTGAT	4860

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•	TGTCCGCAAG	ATGTTATCGC	TGCTGTTAAT	CTGCTGAAGC	GTAAATGCGG	TAATAGGGAC	5100
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	ATTACTTCTG	GTTTGCAGGT	CACTATTGAT	CCGAACAGAA	CGTATCTTAC	GATGACTCAA	5220
	GCTGATAAAG	CGGCCCTTCA	AACGAGGGCT	AAGGATTTTC	CCGTGAGCAA	GGACTGGATT	5280
455	GATGGACACA	TAAAAACAGT	' ACACGAAGCG	CAAGGGATCT	CTGTTGACAA	CGTCACTTTG	5340
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485	CGTCGTGACT	GGGAAAACCC	TGGCGTTACC	CAACTTAATC	GCCTTGCAGC	ACATCCCCCT	6240
	TTCGCCAGCT	GGCGTAATAG	CGAAGAGGCC	CGCACCGATC	GCCCTTCCCA	ACAGTTGCGC	6300
490	AGCCTGAATG	GCGAATGNNN	NNNNAATTCA	GTACATTAAA	AACGTCCGCA	ATGTGTTATT	6360
	AAGTTGTCTA	AGCGTCAATT	TGTTTACACC	ACAATATATC	CTGCCACCAG	CCAGCCAACA	6420
	GCTCCCCGAC	CGGCAGCTCG	GCACAAAATC	ACCACTCGAT	ACAGGCAGCC	CATCAGNNNN	6480

495	иииииииии	имимимими	имимимими	имимимими	иииииииии	иииииииии	6540
	инииииии	имимимими	ииииииииии	имимимими	иииииииии	иниининин	6600
500	иииииииии	имимимими	ииииииииии	имимимими	ииииииииии	инининини	6660
300	ииииииииии	ииииииииии	имимимими	ииииииииии	иииииииии	иииииииии	6720
	ииииииииии	иииииииии	имимимими				6750
<b>50</b> 5							
	<210> 3 <211> 6426 <212> DNA <213> Brom	e mosaic vi	rus				
510	<400> 3						
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	GACAGAÁCCG	CAACGATTGA	AGGAGCCACT	CAGCCGCGG	TTTCTGGAGT	TTAATGAGCT	180
	AAGCACATAC	GTCAGAAACC	ATTATTGCGC	GTTCAAAAGT	CGCCTAAGGT	CACTATCAGO	240
520	TAGCAAATAT	TTCTTGTCAA	AAATGCTCCA	CTGACGTTCC	ATAAATTCCC	CTCGGTATCC	300
	AATTAGNNNN	имимимими	имимимими	GATCGTTTCG	CATGATTGAA	CAAGATGGAT	360
525	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTGG	AGAGGCTATT	CGGCTATGAC	TGGGCACAAC	420
	AGACAATCGG	CTGCTCTGAT	GCCGCCGTGT	TCCGGCTGTC	AGCGCAGGGG	CGCCCGGTTC	480
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530	TATCGTGGCT	GGCCACGACG	GGCGTTCCTT	GCGCAGCTGT	GCTCGACGTT	GTCACTGAAG	600
	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGCA	GGATCTCCTG	TCATCTCACC	660
535	TTGCTCCTGC	CGAGAAAGTA	TCCATCATGG	CTGATGCAAT	GCGGCGGCTG	CATACGCTTG	720
<i>333</i>	ATCCGGCTAC	CTGCCCATTC	GACCACCAAG	CGAAACATCG	CATCGAGCGA	GCACGTACTC	780
•	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	AGAGCATCAG	GGGCTCGCGC	840
540	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	CGGCGATGAT	CTCGTCGTGA	900
	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTCA	960
545	TCGACTGTGG (	CCGGCTGGGT (	GTGGCGGACC (	GCTATCAGGA	CATAGCGTTG	GCTACCCGTG	1020
545	ATATTGCTGA	AGAGCTTGGC (	GCGAATGGG (	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG	1080
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550	мимимими	пимимими в	GATCGTTCAA	ACATTTGGCA	ATAAAGTTTC	TTAAGATTGA	1200
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560	TCTGGTGGTG	GTTCTGGTGG	CGGCTCTGAG	GGTGGTGGCT	CTGAGGGTGG	CGGTTCTGAG	1500
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565	GAAAAGATGG	CAAACGCTAA	TAAGGGGGCT	ATGACCGAAA	ATGCCGATGA	AAACGCGCTA	1620
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	TTCCGTCAAT	ATTTACCTTC	CCTCCCTCAA	TCGGTTGAAT	GTCGCCCTTT	TGTCTTTGGC	1860
575	CCAATACGCA	AACCGCCTCT	CCCCGCGCGT	TGGCCGATTC	ATTAATGCAG	CTGGCACGAC	1920
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590	TAGCCAAAGT	GGTCTGCCTG	ACCAGGAGTT	TTTAACCTTA	ACCAAAGGGC	TGTTCACAGC	2400
	TTAGGTTCAT	ATATCATAGA	ACCGATCATC	TCAGATCAGA	GGGCTTAAAA	GTCTCACAAT	2460
595	GGGACTTCAC	GAGCAAAGCA	TCAACTGACG	TTAGGCCTCC	TCTACCGGTA	GCGTAATCGT	2520
	CGACCTTCTT	TTTCAAGCGT	TGTGTGGTCC	TACGATCATT	AGCTAATTTG	AGTGACTCAC	2580
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	TOTAL COMM	CTCD B B CCDC	TO A CO A TO A CO	marran acoma	maccamacana	COMMUNICA NOM	2264

605	IACATACAG	G AICGCICATO	C TGATAAACT	2 TGATGCCTTC	2 GGTACAGTAG	CAATCAGAGA	282
003	ACCTCAGGA	A ATTCTCGGA	LAADAAATAT E	A AAGCCGCAAG	G AGCAGCTCTA	ACCTCCTCGA	288
	AAATCCAAG	G TTTTTCTTT	C CCATATTTC	GATAAACAA	A ATGACAGAGO	GTCGTAATCA	294
610	TCTTCTCAT	C AAGTTGATT	A ATAAACTTC	TTCGATCAC	GAAGGAAACG	AAATGTGCTC	300
	TGAGCATCT	G TTCATCACGO	C AGAATCTTTC	GCTTAGCTA	GCGCTGGATC	TCTCTCAGAG	306
615	GATCTGGTA	C AGACACCAA	A TTGCCCATTI	CAGTTTCGAC	GAGAAACTTA	CTACAAACGT	312
	AGGGCACAC	I' AGGGTCCATC	ACTTITATCI	CCATATTGA	GAGAGACGTA	AACATATCGG	318
	TATCCAGGA	TGGCTTAACT	TTAGAGATGA	TTAAAGAATC	ATCTCCTGAA	AATATTGCAC	324
620	AGTCACAGT	C ACTTAGATCA	GAGGCATATG	CAATCATAGO	CATAGTGACA	AGAGTATTAC	3300
	CGAAATATGT	T AAACGCGTCA	CCAGTTCTGC	GTTGGAAGGA	AACGGACATT	CCCACCTTGG	3360
625	CATGAGGGT	TGATAAATAA	GAATCGCGAT	GAAAATCAGA	CCACCAATTC	GTCAGCGGCG	3420
	CTGGAAAGCC	CAGCGCAAGG	AGTATCTCTC	TCTGAAACTC	TAGGTGCAGC	TCACCCTGAG	3480
	ATTTATCAAA	TTTGCTTAGG	TCCGCTTCAA	GAAAGTATCT	GTTATTCAAG	CGGACATTCT	3540
630	TAAGCTCCAG	AGAGGATATC	TTTCCGATAG	GCACAATGAA	CCTGGATTTC	AGGGCCAGTG	3600
	ATAACTTCTC	GAAACAAGCA	GTGAAAAAGG	GTGAAAAATT	ACTAGTCACA	CCTTTACTAT	3660
635	GAAATGTTAT	AGTAGCTGCT	ACTGCTCGTT	CCAAGTGAAG	GGTGTCAGTT	ACAACAGGTT	3720
	TTACGTCAGA	CTTCAGCATA	TGCTGGTACC	GACATAAATC	AGTCTCTGCT	GCCACATTCA	3780
	CACCTTGCAA	GTCCATGTGC	TTACCCCACT	TCTTATGGTA	CTCAAGACAT	TTAGTCATGA	3840
640	CATCCATAGA	AGCTCTCAGA	CAGTCTTCAC	CGTCAACATT	AAGGAATGTG	CTACGAAAGC	3900
	GCTTTGCTAT	AGCTTTCGCA	GTGTCCTTCA	TGTTAATCGC	GTCTCCCATT	TCTGGAACGT	3960
645	CCGCGTTTCG	CTTTTTGAGT	GCGGTTAAGA	CTTCTTTCTG	AGTACCAACT	CTTCGCTGAG	4020
	CACTCCCGAT	ATTCATTTTT	GGTTGAAAAT	ATTTATCGGG	GTCCCTATAC	CAGTCTACAT	4080
	CACTTTGCTT	AAGTCTGATC	CTATCAAAGT	CCATGGAATA	ATCACCATTT	TCAACAAGGG	4140
650	CTTGATGGTA	CGAATCATCG	AAATAAGCAT	GGGTTGGCAG	TATGGAATGA	CTGGTCGCTT	4200
	CTGTTCTAGC	AAGGCTGACT	CTCTCCATAT	AAATTGGCCC	AGTAGAGATG	TCAGGGTTAT	4260
655	CTGGATGGCA	GTGTGTATCA	ATAACACGCG	AAACCCTATG	TTCAATAGGG	TTCATGATTT	4320
	GAAGAGTGAT	GTCGTAATCA	GTATTAGTAG	TCTGAAACTC	TTCATCAATG	CCCATGTACC	4380

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PCT/US99/11250

	INICICCAN	G GGTCAGCTCC	TIGGGGGTA:	r CTCCAGTAA	CACGAACTTCC	TCAATTTCAC	444
660	AGTTCGAGG.	A ATCACTGGCC	G AGTTTTAGAT	CGCTCGCAT	G ATCTTCATCG	GCGGCAAACG	450
	ATACACCGT	A ACCATCACTA	GTATCCTCG	GATACCAGTO	ATCAATTTCA	TCTTCGAGCA	456
665	CGAAAGAGC	C CGGAATGTC	AGATATAACA	A TCCGTGCCAT	TTCAGCTTGA	GGAATCAGCG	462
	GTCTATCGG	r gaactgttga	ACCATTTGTT	GGACGGTGTC	GCAAATAGAG	CCCCAGCGCA	468
	CTCGGTCAA	A AGGGGGATCG	AATACCCCTC	CTATCTCCA	GGGCGCTATA	GCTAATTTAA	4740
670	AACTCGCGAC	G AGATCCGTCA	ATGGCAACTC	CGTCTGCCG	CTCCTGCACC	TGAAGGCTAG	4800
	CAGCCTCCAC	CTCGTCTTCT	AAGGATTGAT	CTATGATCC	TTGGAAAGAC	GGGACCTGGC	4860
675	GAACGAAATO	ATCATCCCAG	GTTTTCGAAG	ACATCTTGGT	GATAGTAGAA	AGAACAAGCA	4920
	CACAACAACA	A ACAAGGTCAG	ATGTGTGTTG	CGGGTACCGA	GCTCGAATTC	TCGAGGTCCT	4980
	CTCCAAATGA	AATGAACTTC	CTTATATAGA	GGAAGGGTCT	TGCGAAGGAT	AGTGGGATTG	5040
680	TGCGTCATCC	CTTACGTCAG	TGGAGATATC	ACATCAATCC	ACTTGCTTTG	AAGACGTGGT	5100
	TGGAACGTCT	TCTTTTTCCA	CGATGTTCCT	CGTGGGTGGG	GGTCCATCTT	TGGGACCACT	5160
685	GTCGGTAGAG	GCATTCTTGA	ACGATAGCCT	TTCCTTTATC	GCAATGATGG	CATTTGTAGA	5220
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690	AGACTGTATC	TTTGATATTC	TTGGAGTAGA	CGAGAGTGTC	GTGCTCCACC	ATGTTGAĆCT	5400
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695	CTTATGTTAC	GTCCTGTAGA	AACCCCAACC	CGTGAAATCA	AAAAACTCGA	CGGCCTGTGG	5520
	GCATTCAGTC	TGGATCGCGA	AAACTGTGGA	ATTGATCAGC	GTTGGTGGGA	AAGCGCGTTA	5580
	CAAGAAAGCC	GGGCAATTGC	TGTGCCAGGC	AGTTTTAACG	ATCAGTTCGC	CGATGCAGAT	5640
700	ATTCGTAATT	ATGCGGGCAA	CGTCTGGTAT	CAGCGCGAAG	TCTTTATACC	GAAAGGTTGG	5700
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,03	CCGTATGTTA	TTGCCGGGAA	AAGTGTACAA	TTCACTGGCC	GTCGTTTTAC	AACGTCGTGA	5880
	CTGGGAAAAC	CCTGGCGTTA	CCCAACTTAA	TCGCCTTGCA	GCACATCCCC	CTTTCGCCAG	5940
710	CTGGCGTAAT	AGCGAAGAGG	CCCGCACCGA	TCGCCCTTCC	CAACAGTTGC	GCAGCCTGAA	6000

	TGGCGAATGI	ТТААИИИИИ И	CAGTACATTA	AAAACGTCCG	CAATGTGTTA	TTAAGTTGTC	6060
715	TAAGCGTCAA	TTTGTTTACA	CCACAATATA	TCCTGCCACC	AGCCAGCCAA	CAGCTCCCCG	6120
715	ACCGGCAGCT	CGGCACAAA	TCACCACTCG	ATACAGGCAG	CCCATCAGNN	инининини	6180
	инининини	имимимими	ииииииииии	иииииииии	иииииииии	иииииииииии	6240
720	иииииииии	<b>นทุนหนุนทุนหนุ</b>	ииииииииии	иииииииии	иниининини	имимимими	6300
	имимимими	NNNNNNNNNN	ทททททุททท	имимимими	ииииииииии	ииииииииии	6360
725	имимимими	имимимим	ииииииииии	имимимими	ииииииииии	ииииииииии	6420
125	NNNNNN						6426
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	<212> DNA <213> Brom	e mosaic vi	rus				
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735	AAACACTGAT	' AGTTTAAACT	GAAGGCGGGA	AACGACAATC	TGATCATGAG	CGGAGAATTA	60
	AGGGAGTCAC	GTTATGACCC	CCGCCGATGA	CGCGGGACAA	GCCGTTTTAC	GTTTGGAACT	120
740	GACAGAACCG	CAACGATTGA	AGGAGCCACT	CAGCCGCGGG	TTTCTGGAGT	TTAATGAGCT	180
	AAGCACATAC	GTCAGAAACC	ATTATTGCGC	GTTCAAAAGT	CGCCTAAGGT	CACTATCAGC	240
	TAGCAAATAT	TTCTTGTCAA	AAATGCTCCA	CTGACGTTCC	ATAAATTCCC	CTCGGTATCC	300
745	AATTAGNNNN	имимимими	ииииииииии	GATCGTTTCG	CATGATTGAA	CAAGATGGAT	360
	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTGG	AGAGGCTATT	CGGCTATGAC	TGGGCACAAC	420
750	AGACAATCGG	CTGCTCTGAT	GCCGCCGTGT	TCCGGCTGTC	AGCGCAGGGG	CGCCCGGTTC	480
.20	TTTTTGTCAA	GACCGACCTG	TCCGGTGCCC	TGAATGAACT	GCAGGACGAG	GCAGCGCGGC	540
	TATCGTGGCT	GGCCACGACG	GGCGTTCCTT	GCGCAGCTGT	GCTCGACGTT	GTCACTGAAG	600
755	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGCA	GGATCTCCTG	TCATCTCACC	660
	TTGCTCCTGC	CGAGAAAGTA	TCCATCATGG	CTGATGCAAT	GCGGCGGCTG	CATACGCTTG	720
760	ATCCGGCTAC	CTGCCCATTC	GACCACCAAG	CGAAACATCG	CATCGAGCGA	GCACGTACTC	780
700	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	AGAGCATCAG	GGGCTCGCGC	840
	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	CGGCGATGAT	CTCGTCGTGA	900
765	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTCA	960

	TCGACTGTGG	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	CATAGCGTTG	GCTACCCGTG	102
770	ATATTGCTGA	AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG	1086
,,,	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	TGACGAGTTC	TTCTGANNNN	1140
	иииииииии	инииминии	GATCGTTCAA	ACATTTGGCA	ATAAAGTTTC	TTAAGATTGA	1200
775 ·	ATCCTGTTGC	CGGTCTTGCG	ATGATTATCA	TATAATTTCT	GTTGAATTAC	GTTAAGCATG	1260
	TAATAATTAA	CATGTAATGC	ATGACGTTAT	TTATGAGATG	GGTTTTTATG	ATTAGAGTCC	1320
780	CGCAATTATA	CATTTAATAC	GCGATAGAAA	ACAAAATATA	GCGCGCAAAC	TAGGATAAAT	1,380
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	GAAAAGATGG	CAAACGCTAA	TAAGGGGGCT	ATGACCGAAA	ATGCCGATGA	AAACGCGCTA	1620
790	CAGTCTGACG	CTAAAGGCAA	ACTTGATTCT	GTCGCTACTG	ATTACGGTGC	TGCTATCGAT	1680
	GGTTTCATTG	GTGACGTTTC	CGGCCTTGCT	AATGGTAATG	GTGCTACTGG	TGATTTTGCT	1740
	GGCTCTAATT	CCCAAATGGC	TCAAGTCGGT	GACGGTGATA	ATTCACCTTT	AATGAATAAT	1800
795	TTCCGTCAAT	ATTTACCTTC	CCTCCCTCAA	TCGGTTGAAT	GTCGCCCTTT	TGTCTTTGGC	1860
	CCAATACGCA	AACCGCCTCT	CCCCGCGCGT	TGGCCGATTC	ATTAATGCAG	CTGGCACGAC	1920
800	AGGTTTCCCG	ACTGGAAAGC	GGGCAGTGAG	CGCAACGCAA	TTAATGTGAG	TTAGCTCACT	1980
	CATTAGGCAC	CCCAGGCTTT	ACACTTTATG	CTTCCGGCTC	GTATGTTGTG	TGGAATTGTG	2040
	AGCGGATAAC	AATTTCACAC	AGGAAACAGC	TATGACCATG	ATTACGCCAA	GCTTGCTGCC	2100
805	TGCAGGTCAA	CATGGTGGAG	CACGACACTC	TCGTCTACTC	CAAGAATATC	AAAGATACAG	2160
	TCTCAGAAGA	CCAGAGGGCT	ATTGAGACTT	TTCAACAAAG	GGTAATATCG	GGAAACCTCC	2220
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	GCTTCTACAA	ATGCCATCAT	TGCGATAAAG	GAAAGGCTAT	CGTTCAAGAA	TGCCTCTACC	2340
	GACAGTGGTC	CCAAAGATGG	ACCCCCACCC	ACGAGGAACA	TCGTGGAAAA	AGAAGACGTT	2400
815	CCAACCACGT	CTTCAAAGCA	AGTGGATTGA	TGTGATATCT	CCACTGACGT	AAGGGATGAC	2460
	GCACAATCCC	ACTATCCTTC	GCAAGACCCT	TCCTCTATAT	AAGGAAGTTC	ATTTCATTTG	2520
820	GAGAGGACCT	CGAGAATTCG	AGCTCGGTAC	CCGCAACACA	CATCTGACCT	TGTTGTTGTT	2580

	GTGTGCTTGT	TCTTTCTACT	ATCACCAAGA	TGTCTTCGAA	AACCTGGGAT	GATGATTTCG	2640
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	GTTTTAAATT	AGCTATAGCG	CCCTTGGAGA	TAGGAGGGGT	ATTCGATCCC	CCTTTTGACC	2820
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•	TCGTGCTCGA	AGATGAAATT	GATGACTGGT	ATCCCGAGGA	TACTAGTGAT	GGTTACGGTG	3000
835	TATCGTTTGC	CGCCGATGAA	GATCATGCGA	GCGATCTAAA	ACTCGCCAGT	GATTCCTCGA	3060
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845	CAGAAGCGAC	CAGTCATTCC	ATACTGCCAA	CCCATGCTTA	TTTCGATGAT	TCGTACCATC	3360
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855	AGCGCTTTCG	TAGCACATTC	CTTAATGTTG	ACGGTGAAGA	CTGTCTGAGA	GCTTCTATGG	3660
	ATGTCATGAC	TAAATGTCTT	GAGTACCATA	AGAAGTGGGG	TAAGCACATG	GACTTGCAAG	3720
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865	TATCACTGGC	CCTGAAATCC	AGGTTCATTG	TGCCTATCGG	AAAGATATCC	TCTCTGGAGC	3960
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	CAGCGCCGCT	GACGAATTGG	TGGTCTGATT	TTCATCGCGA	TTCTTATTTA	TCAGACCCTC	4140
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	ACTGTGCAAT	ATTTTCAGGA	GATGATTCTT	TAATCATCTC	TAAAGTTAAG	CCAGTCCTGG	4320
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			, ,			
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935	TGGCGAATGN NNNNNNA	ATT CAGTACATT	AAAACGTCCG	CAATGTGTTA	TTAAGTTGTC	6060
	TAAGCGTCAA TTTGTTT	ACA CCACAATATA	TCCTGCCACC	AGCCAGCCAA	CAGCTCCCCG	6120
940	ACCGGCAGCT CGGCACA	AAA TCACCACTCO	ATACAGGCAG	CCCATCAGNN	имимимими	6180
	имимими имимимими	иимимими им	имимимими	ииииииииии	имимимими	6240
	имимими мимимими	инининини ин	<b>תתתתתתתתת</b>	имимимими	имимимими	6300
<del>9</del> 45	инипипин инипипини	<b>กนนนนนนนน นน่</b> เ	инининини	ииииииииии	ииииииииии	6360
	ининини инининини	NUNUNUNUNU NUI	имимимими	имимимими	имимимими	6420
950	ทุกทุกทุกทุก ทุกทุกทุกทุก	имимимими им	ииииииииии	иииииииии	ииииииииии	6480
930	имимими имимимими	ии				6500
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	TAGCAAATAT TTCTTGTC	AA AAATGCTCC	CTGACGTTCC	ATAAATTCCC	CTCGGTATCC	300
970	ААТТАСИНИИ ИНИИИНИ	แนนหนนหนน นน	GATCGTTTCG	CATGATTGAA	CAAGATGGAT	360
	TGCACGCAGG TTCTCCGG	CC GCTTGGGTGG	AGAGGCTÁTT	CGGCTATGAC	TGGGCACAAC	420
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7.75	TTTTTGTCAA GACCGACC	TG TCCGGTGCCC	TGAATGAACT	GCAGGACGAG	GCAGCGCGGC	540
	TATCGTGGCT GGCCACGA	CG GGCGTTCCT1	GCGCAGCTGT	GCTCGACGTT	GTCACTGAAG	600
980	CGGGAAGGGA CTGGCTGC	TA TTGGGCGAAG	TGCCGGGGCA	GGATCTCCTG	TCATCTCACC	660
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985	ATCCGGCTA	C CTGCCCATT	GACCACCAA	G CGAAACATC	G CATCGAGCG	A GCACGTACT	C 780
	GGATGGAAG	C CGGTCTTGT	C GATCAGGAT	G ATCTGGACG	A AGAGCATCA	3 GGGCTCGCG	C 840
	CAGCCGAAC	T GTTCGCCAG	G CTCAAGGCG	C GCATGCCCG	A CGGCGATGA	r ctcgtcgtg	A 900
990	CCCATGGCG.	A TGCCTGCTTC	CCGAATATC	A TGGTGGAAA	A TGGCCGCTT	r TCTGGATTC	A 960
	TCGACTGTGC	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	CATAGCGTTG	GCTACCCGTG	1020
995	ATATTGCTGA	A AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG	1080
,,,	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	TGACGAGTTC	TTCTGANNNN	1140
	имимимими	นทนทนทนทน เ	GATCGTTCAA	ACATTIGGCA	ATAAAGTTTC	TTAAGATTGA	1200
1000	ATCCTGTTGC	CGGTCTTGCG	ATGATTATCA	TATAATTTCT	GTTGAATTAC	GTTAAGCATG	1260
	TAATAATTAA	CATGTAATGC	ATGACGTTAT	TTATGAGATG	GGTTTTTATG	ATTAGAGTCC	1320
1005	CGCAATTATA	CATTIAATAC	GCGATAGAAA	ACAAAATATA	GCGCGCAAAC	TAGGATAAAT	1380
	TATCGCGCGC	GGTGTCATCT	ATGTTACTAG	ATCGGGCCTC	CTGTCAATGC	TGGCGGCGGC	1440
	TCTGGTGGTG	GTTCTGGTGG	CGGCTCTGAG	GGTGGTGGCT	CTGAGGGTGG	CGGTTCTGAG	1500
1010	GGTGGCGGCT	CTGAGGGAGG	CGGTTCCGGT	GGTGGCTCTG	GTTCCGGTGA	TTTTGATTAT	1560
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CCO	Thearmore	Chohohom	aama s mmmaa	001000000			

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26

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1712							

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1425	ATAACTTCT	C GAAACAAGCA	A GTGAAAAAGG	GTGAAAAATT	CACTAGTCACA	CCTTTACTAT	366
, ,	GAAATGTTA'	T AGTAGCTGCT	ACTGCTCGTT	' CCAAGTGAAG	GGTGTCAGTT	ACAACAGGTT	372
	TTACGTCAG	A CTTCAGCATA	TGCTGGTACC	GACATAAATC	AGTCTCTGCT	GCCACATTCA	378
1430	CACCTTGCA	A GTCCATGTGC	TTACCCCACT	TCTTATGGTA	CTCAAGACAT	TTAGTCATGA	. 3840
	CATCCATAG	A AGCTCTCAGA	CAGTCTTCAC	CGTCAACATT	AAGGAATGTG	CTACGAAAGC	3900
1435	GCTTTGCTAT	r AGCTTTCGCA	GTGTCCTTCA	TGTTAATCGC	GTCTCCCATT	TCTGGAACGT	3960
	CCGCGTTTCC	CTTTTTGAGT	GCGGTTAAGA	CTTCTTTCTG	AGTACCAACT	CTTCGCTGAG	4020
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1440	CACTTTGCTT	AAGTCTGATC	CTATCAAAGT	CCATGGAATA	ATCACCATTT	TCAACAAGGG	4140
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1450	TATCTCCAAG	GGTCAGCTCC	TTGGGGGTAT	CTCCAGTAAC	ACGAACTTCC	TCAATTTCAC	4440
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	CGAAAGAGCC	CGGAATGTCA	AGATATAACA	TCCGTGCCAT	TTCAGCTTGA	GGAATCAGCG	4620
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1460	CTCGGTCAAA	AGGGGGATCG	AATACCCCTC	СТАТСТССАА	GGGCGCTATA	GCTAATTTAA	4740
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	GAACGAAATC	ATCATCCCAG	GTTTTCGAAG	ACATCTTGGT	GATAGTAGAA	AGAACAAGCA	4920
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1470	CTCCAAATG	A AATGAACTIC	CTTATATAG	A GGAAGGGTC	r tgcgaaggai	TAGTGGGATTC	5040
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	GTCGGTAGA	G GCATTCTTGA	ACGATAGCC	TTCCTTTATO	GCAATGATGG	CATTTGTAGA	5220
	AGCCATCTT	C CTTTTCTACT	GTCCTTTCG	A TGAAGTGACA	GATAGCTGGG	CAATGGAATC	5280
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	AGACTGTAT	C TTTGATATTC	TTGGAGTAGA	CGAGAGTGTC	GTGCTCCACC	ATGTTGACCT	5400
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	TGGTTTTAG	G AATTAGAAAT	TTTATTGATA	GAAGTATTTI	· ACAAATACAA	ATACATACTA	5520
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	CTGGTGATT	r GCGGACTCTA	GAGGATCCCC	GGGTACCGAG	CTCGAATTCT	CGAGCAGAGG	5700
1495	TCTCACACAC	3 AGACAAGCGC	ATCACTTAAC	ACAATTAAAG	ATCAAATCAC	CAGCGAGCTC	5760
	GCCGTTAAAG	CAATACTCAA	AGGACTTCTT	GTGTCGTGTT	AAGGCAACCA	AACAGTACTC	5820
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1505	CGTTCTGTTC	GGATCAATAG	TGACCTGCAA	ACCAGAAGTA	ATACGACGCT	TCGTGAGACT	6060
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1510	AACGTCACGG	CGATCATATT	GCAGATTACC	GTGGAGCAAT	TTAAAACCCG	CGTCACGAGA	6240
	CTTGAACGAA	ATCTGCTCTG	TGTCCCCAAA	GGCAAGAACT	TGTGAACATT	TAGACAGAGC	6300
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						ATTTACGATT	
1520	CGCCGTCACA	ATTAGGTCCT	CTCCCATACG	GAATGCATCT	TTTATGGCAG	TGGTTTTACC	6540
	GCATCCCGCA	ACTCCATCAA	CCATGGAAAT	ATCGCATGTA	GGGACAGAAA	CTTTGGCGCT	6600

WO 99/61597

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1525	AGCTTCTGC	A ATGTCCCTCA	AGTTAGAGCA	TGCACATGTT	TTATCAACAA	TGTACGTTTC	666
1323	ATCTGCGTG	C TTCGGACCTA	AACCATGCTC	ATTATATCCA	ACAGTGTAAT	CGTATTTTT	672
	AGGATACAA	C CAGTTACCGT	TGGCCAAATG	GACATTCACC	ATATCGTCTA	TGCGATGGTA	6780
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	GTGACGAAGA	A TTAGACTCGG	AGTTGTTATG	TAATCTCTTA	CAATAACGCA	CAAATTCCTT	6900
1535	CATGGCTCCC	TGTCTAGATA	TGCCAĆGAGG	GTCCGTTGGT	ACCTCAACAG	ACACCTCGGC	6960
	ATCCGGGACC	CACATCAGTCA	CCGGTTTAAC	GTCATCACTG	ACGGACTCAG	GGCTCGAACT	7020
	CTCAGGGGCA	TCATGAAACT	CCTCCTGAGG	TATCTCAGCA	GCTGGCGGGA	CTTTCGCCTT	7080
1540	CTTCTTCGAG	CGCTTGGTCT	TGGCTGTCTG	CACTTCATGC	TCCAGCCGGT	CGAATAAGTC	7140
	CTCTTCAGTC	CAAAACGTTC	TCAAACGTGA	TATCGGTACA	GAATCTTGCT	CAAATTCTTC	7200
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1555	CGACGACTTG	GCGGATAAGA	TAGATGCGAC	AGCTTTCATG	TTCTCAGTCC	ATTCTTTACT	7560
	TTCCTTGAAA	CATCTGAAAG	CTATCTCCTC	TACCTCTCTC	ACTGTGGTTT	TGGCGACGCG	7620
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1560	TATGTCTTCA	AACCATACAC	AGTGACGTAG	TGTCTCCCGG	GGGCAGCGTA	AATTTGTAGC	7740
					CGCTCCAACA		
1565	TCCATCGATG	CAATGCACCG	ACTCGGTGAA	AAATGAGCCC	AAATCTTGCC	ATCCGTGGAT	7860
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1570	GCCGTCGAAC	ATAACGGTAC	CACGTAGTAC	GCGTACTCCA	TGCGAATGCA	TGGCGTCACA	8040
					ATAGCCCAAT		
1575					TCATCGAAAT		
	TTGCAAAATT	TTTCGCATGC	GGCACATCCT	CTCCTCATGT	CGGGCAGCGT	CTCTAACACC	8220

	CANCACAGG	A CHACHACIG	GCACCCTTT	r Arecerrer	r GAAAAGTGAT	GCCACCAAGA	8280
1580	CCCTCCGAA	A TCTATAACGO	GGTCTTCAG	GGGAAAACT	G TCGAGACAGT	CATAATGCTC	8340
	CGCTACACG	C AGAGCACCAC	CCAGGCTATO	G GGGCGCATG	TACTGCTGAG	TCAAATTTAA	8400
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	CCATCTTTGG	GACCACTGTC	GGTAGAGGCA	TTCTTGAACG	ATAGCCTTTC	CTTTATCGCA	8880
1600	ATGATGGCAT	TTGTAGAAGC	CATCTTCCTT	TTCTACTGTC	CTTTCGATGA	AGTGACAGAT	8940
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1605	AGCCCTCTGG	TCTTCTGAGA	CTGTATCTTT	GATATTCTTG	GAGTAGACGA	GAGTGTCGTG	9060
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1615	AGCGCGAAGT	CTTTATACCG	AAAGGTTGGG	CAGGCCAGCG	TATCGTGCTG	CGTTTCGATG	9360
	CGGTCACTCA	TTACGGCAAA	GTGTGGGTCA	ATAATCAGGA	AGTGATGGAG	CATCAGGGCG	9420
	GCTATACGCC	ATTTGAAGCC	GATGTCACGC	CGTATGTTAT	TGCCGGGAAA	AGTGTACAAT	9480
1620	TCACTGGCCG	TCGTTTTACA	ACGTCGTGAC	TGGGAAAACC	CTGGCGTTAC	CCAACTTAAT	9540
	CGCCTTGCAG	CACATCCCCC	TTTCGCCAGC	TGGCGTAATA	GCGAAGAGGC	CCGCACCGAT	9600
1625	CGCCCTTCCC	AACAGTTGCG	CAGCCTGAAT	GGCGAATGNN	NNNNAATTC	AGTACATTAA	9660
1023	AAACGTCCGC	AATGTGTTAT	TAAGTTGTCT	AAGCGTCAAT	TTGTTTACAC	САСААТАТАТ	9720
	CCTGCCACCA	GCCAGCCAAC	AGCTCCCCGA	CCGGCAGCTC	GGCACAAAAT	CACCACTCGA	9780
1630	TACAGGCAGC	CCATCAGNNN	иииииииии	ииииииииии	имимимими	ииииииииии	9840

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1635	иииииииии	имимимими г	имимимими	ииииииииии	иииииииии	иниининии	9960
	NNNNNNNN	นทุนทุนทุนทุน	имимимими	имимимими	имимимими	имимимими	10020
	инининини	имимимими в	имимимими	инининини	иииииииииииииии	имимимими	10080
1640	инининини	имимимими и	ииииииииии	иииииииии	ииииииииии	имимимими	10140
	имимимими	имимимими и	ииииииииии	ииииииииии	имимимими	иииииииии	10200
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1650		ne mosaic vi	rus				
	<400> 7 AAACAÇTGAT	AGTTTAAACT	GAAGGCGGGA	AACGACAATO	TGATCATGA	G CGGAGAATT	A 60
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	GACAGAACCG	CAACGATTGA	AGGAGCCACT	CAGCCGCGG	TTTCTGGAG	r TTAATGAGC	Г 180
	AAGCACATAC	GTCAGAAACC	ATTATTGCGC	GTTCAAAAGT	CGCCTAAGG	CACTATCAG	240
1660	TAGCAAATAT	TTCTTGTCAA	AAATGCTCCA	CTGACGTTCC	: ATAAATTCC	CTCGGTATC	300
	AATTAGNNNN	имимимими	ииииииииии	GATCGTTTCG	CATGATTGA	A CAAGATGGAT	r 360
1665	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTGG	AGAGGCTATT	CGGCTATGAC	TGGGCACAAC	420
	AGACAATCGG	CTGCTCTGAT	GCCGCCGTGT	TCCGGCTGTC	AGCGCAGGGC	CGCCCGGTTC	2 480
	TTTTTGTCAA	GACCGACCTG	TCCGGTGCCC	TGAATGAACT	GCAGGACGAG	GCAGCGCGG	540
1670	TATCGTGGCT	GGCCACGACG	GGCGTTCCTT	GCGCAGCTGT	GCTCGACGTT	GTCACTGAAG	600
	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGCA	GGATCTCCTG	TCATCTCACC	660
1675	TTGCTCCTGC	CGAGAAAGTA	TCCATCATGG	CTGATGCAAT	GCGGCGGCTG	CATACGCTTG	720
	ATCCGGCTAC	CTGCCCATTC	GACCACCAAG	CGAAACATCG	CATCGAGCGA	GCACGTACTC	780
	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	AGAGCATCAG	GGGCTCGCGC	840
1680	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	CGGCGATGAT	CTCGTCGTGA	900
	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTCA	960
1685	TCGACTGTGG	CCGGCTGGGT (	GTGGCGGACC (	<b>ECTATCAGGA</b>	CATAGCGTTG	GCTACCCGTG	1020

	ATATTGCTG	A AGAGCTTGG	GGCGAATGG	G CTGACCGCT	r ccrccrccr	TACGGTATC	3 108
	CCGCTCCCG	A TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTC	TGACGAGTT	TTCTGANNNI	N 114
1690	имимими	<b>ทุกทุกทุกทุก</b> ท	GATCGTTCA	ACATTTGGC	ATANAGTTTO	TTAAGATTG	A 120
	ATCCTGTTG	C CGGTCTTGCG	ATGATTATCA	TATAATTTC	GTTGAATTAC	GTTAAGCATO	3 1260
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	CGCAATTAT.	A CATTTAATAC	GCGATÁGAAA	ACAAAATATA	GCGCGCAAAC	TAGGATAAAT	1380
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1700	TCTGGTGGT	G GTTCTGGTGG	CGGCTCTGAG	GGTGGTGGCT	CTGAGGGTGG	CGGTTCTGAG	1500
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1733	GAGAGGACCT	CGAGAATTCG .	AGCTCGGTAC	CCGCAACACA	CATCTGACCT	TGTTGTTGTT	2580
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33

1740 TTCGCCAGGT CCCGTCTTTC CAATGGATCA TAGATCAATC CTTAGAAGAC GAGGTGGAGG 2700 CTGCTAGCCT TCAGGTGCAG GAGCCGGCAG ACGGAGTTGC CATTGACGGA TCTCTCGCGA 2760 GTTTTAAATT AGCTATAGCG CCCTTGGAGA TAGGAGGGGT ATTCGATCCC CCTTTTGACC 2820 1745 GAGTGCGCTG GGGCTCTATT TGCGACACCG TCCAACAAAT GGTTCAACAG TTCACCGATA 2880 GACCGCTGAT TCCTCAAGCT GAAATGGCAC GGATGTTATA TCTTGACATT CCGGGCTCTT 2940 1750 TCGTGCTCGA AGATGAAATT GATGACTGGT ATCCCGAGGA TACTAGTGAT GGTTACGGTG 3000 TATCGTTTGC CGCCGATGAA GATCATGCGA GCGATCTAAA ACTCGCCAGT GATTCCTCGA 3060 ACTGTGAAAT TGAGGAAGTT CGTGTTACTG GAGATACCCC CAAGGAGCTG ACCCTTGGAG 3120 1755 ATAGGTACAT GGGCATTGAT GAAGAGTTTC AGACTACTAA TACTGATTAC GACATCACTC 3180 TTCAAATCAT GAACCCTATT GAACATAGGG TTTCGCGTGT TATTGATACA CACTGCCATC 3240 1760 CAGATAACCC TGACATCTCT ACTGGGCCAA TTTATATGGA GAGAGTCAGC CTTGCTAGAA 3300 CAGAAGCGAC CAGTCATTCC ATACTGCCAA CCCATGCTTA TTTCGATGAT TCGTACCATC 3360 AAGCCCTTGT TGAAAATGGT GATTATTCCA TGGACTTTGA TAGGATCAGA CTTAAGCAAA 3420 1765 GTGATGTAGA CTGGTATAGG GACCCCGATA AATATTTTCA ACCAAAAATG AATATCGGGA 3480 GTGCTCAGCG AAGAGTTGGT ACTCAGAAAG AAGTCTTAAC CGCACTCAAA AAGCGAAACG 3540 1770 CGGACGTTCC AGAAATGGGA GACGCGATTA ACATGAAGGA CACTGCGAAA GCTATAGCAA 3600 AGCGCTTTCG TAGCACATTC CTTAATGTTG ACGGTGAAGA CTGTCTGAGA GCTTCTATGG 3660 ATGTCATGAC TAAATGTCTT GAGTACCATA AGAAGTGGGG TAAGCACATG GACTTGCAAG 3720 1775 GTGTGAATGT GGCAGCAGAG ACTGATTTAT GTCGGTACCA GCATATGCTG AAGTCTGACG 3780 TAAAACCTGT TGTAACTGAC ACCCTTCACT TGGAACGAGC AGTAGCAGCT ACTATAACAT 3840 1780 TTCATAGTAA AGGTGTGACT AGTAATTTTT CACCCTTTTT CACTGCTTGT TTCGAGAAGT 3900 TATCACTGGC CCTGAAATCC AGGTTCATTG TGCCTATCGG AAAGATATCC TCTCTGGAGC 3960 TTAAGAATGT CCGCTTGAAT AACAGATACT TTCTTGAAGC GGACCTAAGC AAATTTGATA 4020 1785 AATCTCAGGG TGAGCTGCAC CTAGAGTTTC AGAGAGAGAT ACTCCTTGCG CTGGGCTTTC 4080 CAGCGCCGCT GACGAATTGG TGGTCTGATT TTCATCGCGA TTCTTATTTA TCAGACCCTC 4140 1790 ATGCCAAGGT GGGAATGTCC GTTTCCTTCC AACGCAGAAC TGGTGACGCG TTTACATATT 4200 TCGGTAATAC TCTTGTCACT ATGGCTATGA TTGCATATGC CTCTGATCTA AGTGACTGTG 4260

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	ATACCGATA	T GTTTACGTCT	CTCTTCAATA	TGGAGATAA	AGTCATGGAC	CCTAGTGTGC	4380
	CCTACGTTT	G TAGTAAGTTT	CTCGTCGAAA	CTGAAATGG	CAATTTGGTG	TCTGTACCAG	4440
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	TCAGAGCAC	A TTTCGTTTCC	TTCTGTGATC	GAATGAAGTT	TATTAATCAA	CTTGATGAGA	4560
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	GGTTCTGCT	A CTTGTTCTT1	GTTTTTCACC	AACAAAATGT	CAAGTTCTAT	CGATTTGCTG	5940
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	GTTGCGCAA	C AGTTATCTGC	GCAGATTGAA	TACGCGAAAA	GGTCTAAGAA	AATCAACGTT	6060
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	CTGTGTGACG	CCATGCATTC	GCATGGAGTA	CGCGTACTAC	GTGGTACCGT	TATGTTCGAC	6540
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1895	CCCCTACTAC	GTCTGGATTC	TTATGCGGAC	AGTTTTAAGT	TTCTGACTCG	TCTCTCAAAC	7320
	GTTGAAGAAT	TTGAGCAAGA	TTCTGTACCG	ATATCACGTT	TGAGAACGTT	TTGGACTGAA	7380
	GAGGACTTAT	TCGACCGGCT	GGAGCATGAA	GTGCAGACAG	CCAAGACCAA	GCGCTCGAAG	7440
1900	AAGAAGGCGA	AAGTCCCGCC	AGCTGCTGAG	ATACCTCAGG	AGGAGTTTCA	TGATGCCCCT	7500

36

GAGAGTTCGA GCCCTGAGTC CGTCAGTGAT GACGTTAAAC CGGTGACTGA TGTGGTCCCG 7560 GATGCCGAGG TGTCTGTTGA GGTACCAACG GACCCTCGTG GCATATCTAG ACACGGAGCC 7620 1905 ATGAAGGAAT TTGTGCGTTA TTGTAAGAGA TTACATAACA ACTCCGAGTC TAATCTTCGT 7680 CACCTATGGG ACATTTCCGG CGGTCGCGGA AGTGAGATCG CAAATAAGAG CATCTTTGAG 7740 1910 ACCTACCATC GCATAGACGA TATGGTGAAT GTCCATTTGG CCAACGGTAA CTGGTTGTAT 7800 CCTAAAAAAT ACGATTACAC TGTTGGATAT AATGAGCATG GTTTAGGTCC GAAGCACGCA 7860 GATGAAACGT ACATTGTTGA TAAAACATGT GCATGCTCTA ACTTGAGGGA CATTGCAGAA 7920 1915 GCTAGCGCCA AAGTTTCTGT CCCTACATGC GATATTTCCA TGGTTGATGG AGTTGCGGGA 7980 TGCGGTAAAA CCACTGCCAT AAAAGATGCA TTCCGTATGG GAGAGGACCT AATTGTGACG 8040 1920 GCGAATCGTA AATCGGCCGA GGACGTCAGG ATGGCTTTAT TCCCTGACAC TTATAATTCC 8100 AAGGTAGCTT TGGACGTTGT GCGCACCGCG GATTCTGCGA TCATGCACGG TGTACCGTCC 8160 TGTCATAGGC TGCTTGTTGA TGAGGCTGGT TTACTACATT ATGGTCAACT CCTGGTGGTG 8220 1925 GCTGCTCTGT CTAAATGTTC ACAAGTTCTT GCCTTTGGGG ACACAGAGCA GATTTCGTTC 8280 AAGTCTCGTG ACGCGGGTTT TAAATTGCTC CACGGTAATC TGCAATATGA TCGCCGTGAC 8340 1930 GTTGTTCACA AGACTTACCG GTGTCCGCAA GATGTTATCG CTGCTGTTAA TCTGCTGAAG 8400 CGTAAATGCG GTAATAGGGA CACGAAGTAT CAATCCTGGA CATCTGAGTC CAAAGTTTCT 8460 AGAAGTCTCA CGAAGCGTCG TATTACTTCT GGTTTGCAGG TCACTATTGA TCCGAACAGA 8520 1935 ACGTATCTTA CGATGACTCA AGCTGATAAA GCGGCCCTTC AAACGAGGGC TAAGGATTTT 8580 CCCGTGAGCA AGGACTGGAT TGATGGACAC ATAAAAACAG TACACGAAGC GCAAGGGATC 8640 1940 TCTGTTGACA ACGTCACTTT GGTTCGGCTT AAGTCGACCA AATGTGATTT GTTTAAACAT 8700 GAGGAGTACT GTTTGGTTGC CTTAACACGA CACAAGAAGT CCTTTGAGTA TTGCTTTAAC 8760 GGCGAGCTCG CTGGTGATTT GATCTTTAAT TGTGTTAAGT GATGCGCTTG TCTCTGTGTG 8820 1945 AGACCTCTGC TCGAGAATTC GAGCTCGGTA CCCGGGGATC CTCTAGAGTC CGCAAATCAC 8880 CAGTCTCTCT CTACAAATCT ATCTCTCTCT ATTTTCTCCA GAATAATGTG TGAGTAGTTC 8940 1950 CCAGATAAGG GAATTAGGGT TCTTATAGGG TTTCGCTCAT GTGTTGAGCA TATAAGAAAC 9000 CCTTAGTATG TATTTGTATT TGTAAAATAC TTCTATCAAT AAAATTTCTA ATTCCTAAAA 9060 CCAAAATCCA GTGACCGGGT GGTCAGTCCC TTATGTTACG TCCTGTAGAA ACCCCAACCC 9120 1955

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1960	GTTTTAACG	A TCAGTTCGCC	GATGCAGATA	TTCGTAATTA	TGCGGGCAAC	GTCTGGTATC	9300
	AGCGCGAAG:	r CTTTATACCG	AAAGGTTGGG	CAGGCCAGCG	TATCGTGCTG	CGTTTCGATG	9360
1965	CGGTCACTC	TTACGGCAAA	GTGTGGGTCA	ATAATCAGGA	AGTGATGGAG	CATCAGGGCG	9420
	GCTATACGCC	CATTTGAAGCC	GATGTCACGC	CGTATGTTAT	TGCCGGGAAA	AGTGTACAAT	9480
	TCACTGGCCC	TCGTTTTACA	ACGTCGTGAC	TGGGAAAACC	CTGGCGTTAC	CCAACTTAAT	9540
1970	CGCCTTGCAG	CACATCCCCC	TTTCGCCAGC	TGGCGTAATA	GCGAAGAGGC	CCGCACCGAT	9600
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1975	AAACGTCCGC	AATGTGTTAT	TAAGTTGTCT	AAGCGTCAAT	TTGTTTACAC	CACAATATAT	9720
	CCTGCCACCA	GCCAGCCAAC	AGCTCCCCGA	CCGGCAGCTC	GGCACAAAAT	CACCACTCGA	9780
	TACAGGCAGC	CCATCAGNNN	ииииииииии	ииииииииии	ииииииииии	имимимими	9840
1980	NNNNNNNNN	иииииииии	имимимими	NNNNNNNN	иииииииииииииииииииииииииииииииииииииии	ииииииииии	9900
	имимимии	иииииииииии	имимимими	NNNNNNNNN	имимимии	ииииииииии	9960
1985.	иииииииии	ииииииииии	имимимими	иммимимии	имимимими	NNNNNNNNN 1	0020
	имимимими	имимимими	имимимими	имимимими	имимимими	NNNNNNNNN 1	0080
	имимимими	имимимими	инининини	инининини	нимимими	NNNNNNNNN 1	0140
1990	NNNNNNNNN	имимимими	имимимими	имимимими	<b>ו אואו</b> אואאאא	NNNNNNNNN 1	0200
	иииииииииии	ииииииииии	имимимими	имимимими	ממאמממממ	ииииииии 1	0260
1995	имимимими	NN				1	0272
2000	<210> 8 <211> 10166 <212> DNA <213> Brome	6 e mosaic vil	· fus	·			
2000	<400> 8	ACCOMPANA A CON	Ch boooggab	11.001.011			Ω
						CGGAGAATTA	
2005						GTTTGGAACT	
						TTAATGAGCT	
	CAINC	GICHGWWWCC	WITH I I GCGC	GIICAAAAGT	CGCCTAAGGT	CACTATCAGC	240

2010	TAGCAAATAT	TTCTTGTCA	AAATGCTCC.	A CTGACGTTC	C ATAAATTCC	CTCGGTATC	C 300
	AATTAGNNNN	иииииииии	ממאממממממ ו	N GATCGTTTC	G CATGATTGA	A CAAGATGGA	Т 360
2015	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTG	G AGAGGCTAT	CGGCTATGA	C TGGGCACAA	C 420
2013	AGACAATCGG	CTGCTCTGAT	GCCGCCGTG	r TCCGGCTGT	C AGCGCAGGG	3 CGCCCGGTT	C 480
	TTTTTGTCAA	GACCGACCTG	TCCGGTGCC	C TGAATGAAC	CAGGACGA	GCAGCGCGG	C 540
2020	TATCGTGGCT	GGCCACGACG	GGCGTTCCT	r gcgcagctgt	GCTCGACGT	r gtcactgaa	G 600
	CGGGAAGGGA	CTGGCTGCTA	TIGGGCGAA	G TGCCGGGGC	GGATCTCCT	TCATCTCAC	C 660
2025	TTGCTCCTGC	CGAGAAAGTA	TCCATCATG	G CTGATGCAAT	GCGGCGGCTG	CATACGCTT	G 720
2023	ATCCGGCTAC	CTGCCCATTC	GACCACCAA	G CGAAACATCO	CATCGAGCGA	GCACGTACT	C 780
	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	S ATCTGGACGA	AGAGCATCAG	GGGCTCGCG	C 840
2030	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	C GCATGCCCGA	CGGCGATGAT	CTCGTCGTG	A 900
	CCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTC	A 960
2035	TCGACTGTGG	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	CATAGCGTTG	GCTACCCGTG	1020
	ATATTGCTGA	AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG	1080
	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	TGACGAGTTC	TTCTGANNNN	1140
2040	NNNNNNNNN	имимимими	GATCGTTCAA	ACATTTGGCA	ATAAAGTTTC	TTAAGATTGA	1200
	ATCCTGTTGC	CGGTCTTGCG	ATGATTATCA	TATAATTTCT	GTTGAATTAC	GTTAAGĆATG	1260
2045	AATTAATAAT	CATGTAATGC	ATGACGTTAT	TTATGAGATG	GGTTTTTATG	ATTAGAGTCC	1320
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	CATTAGGCAC	CCCAGGCTTT	ACACTTTATG	CTTCCGGCTC	GTATGTTGTG	TGGAATTGTG	2040
2070	AGCGGATAAC	AATTTCACAC	AGGAAACAGC	TATGACCATG	ATTACGCCAA	GCTTGCTGCC	2100
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	GCTTCTACAA	ATGCCATCAT	TGCGATAAAG	GAAAGGCTAT	CGTTCAAGAA	TGCCTCTACC	2340
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	CCAACCACGT	CTTCAAAGCA	AGTGGATTGA	TGTGATATCT	CCACTGACGT	AAGGGATGAC	2460
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	GTGTGCTTGT	TCTTTCTACT	ATCACCAAGA	TGTCTTCGAA	AACCTGGGAT	GATGATTTCG	2640
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2110	TTCAAATCAT	GAACCCTATT	GAACATAGGG	TTTCGCGTGT	TATTGATACA	CACTGCCATC	3240
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2210	CTTGAACGAA	ATCTGCTCTG	TGTCCCCAAA	GGCAAGAACT	TGTGAACATT	TAGACAGAGC	6300
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	GGTCTCAAA	G ATGCTCTTAT	TTGCGATCTC	ACTTCCGCGA	CCGCCGGAAA	TGTCCCATAG	6841
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	CGACGACTTG	GCGGATAAGA	TAGATGCGAC	AGCTTTCATG	TTCTCAGTCC	ATTCTTTACT	7560
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# **Claims**

ì	1. A DNA-launching platform comprising:
2	a) a polynucleotide molecule encoding a modified viral RNA molecule; and
3	b) a DNA dependent RNA polymerase promoter.
1	2. The DNA-launching platform of claim 1 further comprising a sequence encoding a
2	least one cis-acting element.
I	3. The DNA-launching platform of claim 1 further comprising a ribozyme sequence
1	4. The DNA-launching platform of claim 1 further comprising a termination sequence
1	5. The DNA-launching platform of claim 1 further comprising a restriction site.
1	6. The DNA-launching platform of claim 1 wherein said modified RNA molecule
2	comprises an exogenous RNA segment.
1	7. The DNA-launching platform of claim I wherein said DNA dependent RNA
2	polymerase promoter is capable of functioning in a plant cell.
ì	8. A method of genotypically or phenotypically modifying one or more cells comprising
2	the following steps:
3	a) obtaining a DNA-launching platform comprising a polynucleotide molecule encoding
4	a modified viral RNA; and
	b) transfecting said one or more cells with said DNA-launching platform, wherein said
	polynucleotide molecule is transcribed thereby forming a replicatable RNA transcript.
ı	9. The method of claim 8 further comprising pre-transforming said cell with at least one
2	polynucleotide molecule encoding at least one trans-acting factor.
1	10. The method of claim 8 further comprising introducing a trans-acting factor.

- ١ 11. The method of claim 10 wherein said introducing a trans-acting factor comprises 2 co-transfection of an expression plasmid comprising a nucleotide sequence encoding said trans-3 acting factor. ı 12. The method of claim 10 wherein said introducing a trans-acting factor comprises 2 co-transfection of an RNA transcript encoding said trans-acting factor. 1 13. The method of claim 10 wherein said trans-acting factor is stably expressed. 14. The method of claim 8 wherein said modified viral RNA comprises an exogenous 2 RNA segment. I 15. The method of claim 8 wherein said DNA-launching platform comprises a ribozyme 2 sequence. 1 The method of claim 8 wherein said DNA-launching platform comprises a 2 promoter. 1 17. The method of claim 8 wherein said DNA-launching platform comprises a 2 termination sequence. 1 18. The method of claim 8 wherein said DNA-launching platform comprises a 2 restriction site. 1 19. The modified cell produced by the method of claim 8. 1 20. A method of producing a plant or plant tissue comprising at least one genotypically 2 or phenotypically modified cell, said method comprising transfecting cells of said plant or plant 3 tissue with a DNA-launching platform, wherein said DNA-launching platform comprises a 4 polynucleotide encoding a modified RNA molecule, such that said polynucleotide molecule is 5 transcribed to form a replicatable RNA transcript. 1 21. The method of claim 20 wherein said modified RNA molecule comprises an
- 2 exogenous RNA segment.

١ 22. The method of claim 20 wherein said DNA-launching platform comprises a 2 ribozyme sequence. ı 23. The method of claim 20 wherein said DNA-launching platform comprises a 2 promoter. 1 24. The method of claim 20 wherein said DNA-launching platform comprises a 2 termination sequence. l 25. The method of claim 20 wherein said DNA-launching platform comprises a 2 restriction site. 1 26. A method of producing a genotypically or phenotypically modified plant comprising 2 obtaining at least one modified cell produced by the method of claim 8; and subjecting said modified cell to conditions whereby a plant is regenerated therefrom. 3 1 27. A plant produced by the method of claim 26. 28. A plant descended from the plant of claim 27. 1 29. The method of claim 20, wherein said plant or plant tissue comprises one or more 2 cells transformed with a polynucleotide molecule encoding at least one trans-acting factor, 3 wherein said polynucleotide molecule is expressed. 30. The method of claim 29, wherein said modified viral RNA molecule is capable of 1 2 replication only in said one or more cells transformed with a polynucleotide molecule encoding 3 at least one trans-acting factor.

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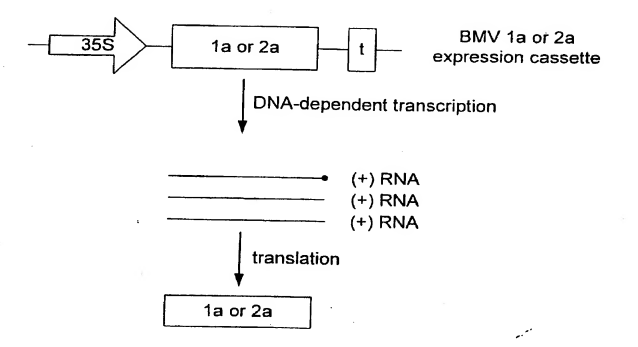


FIG. 1

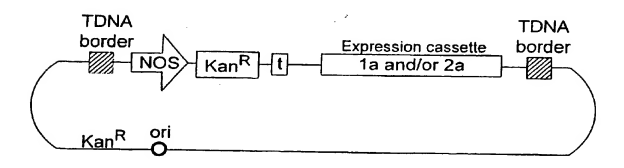


FIG. 2

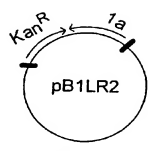
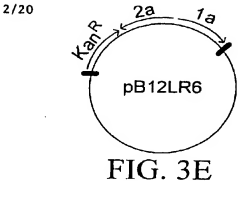


FIG. 3A



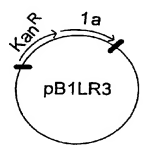
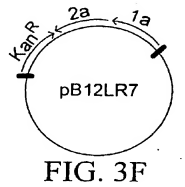
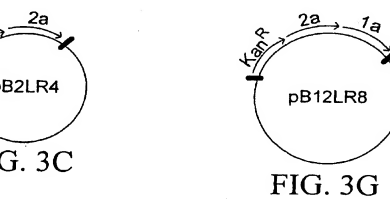
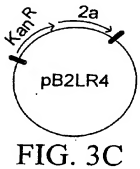
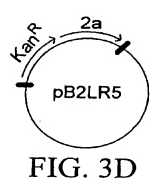


FIG. 3B

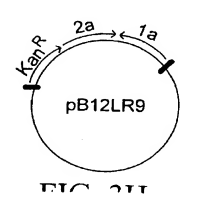








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3/20

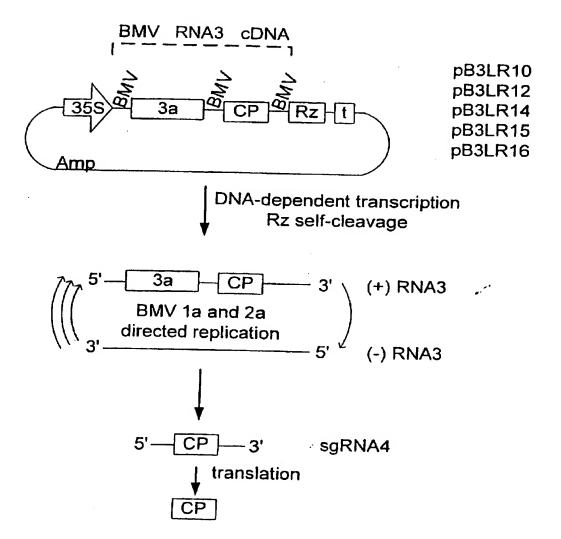


FIG. 4

WO 99/61597

**Amp** 

PCT/US99/11250

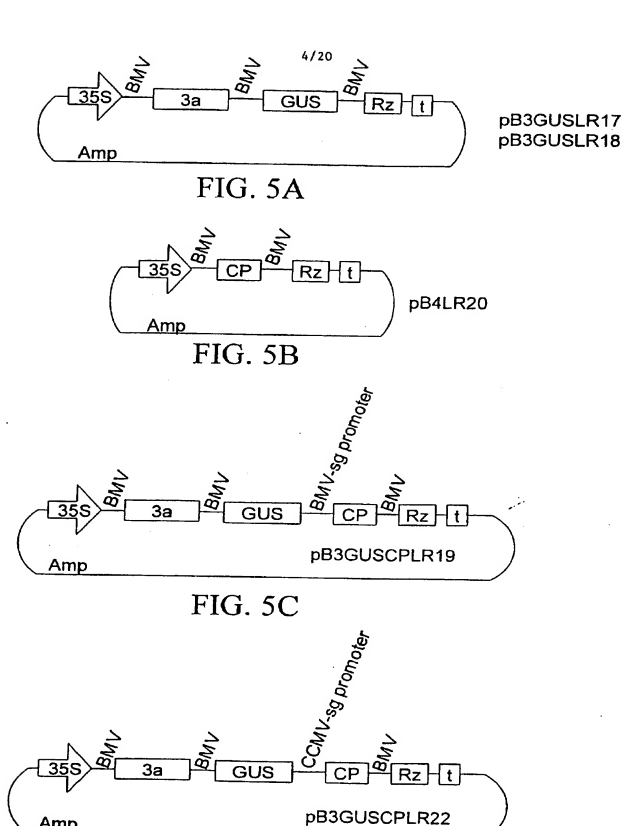
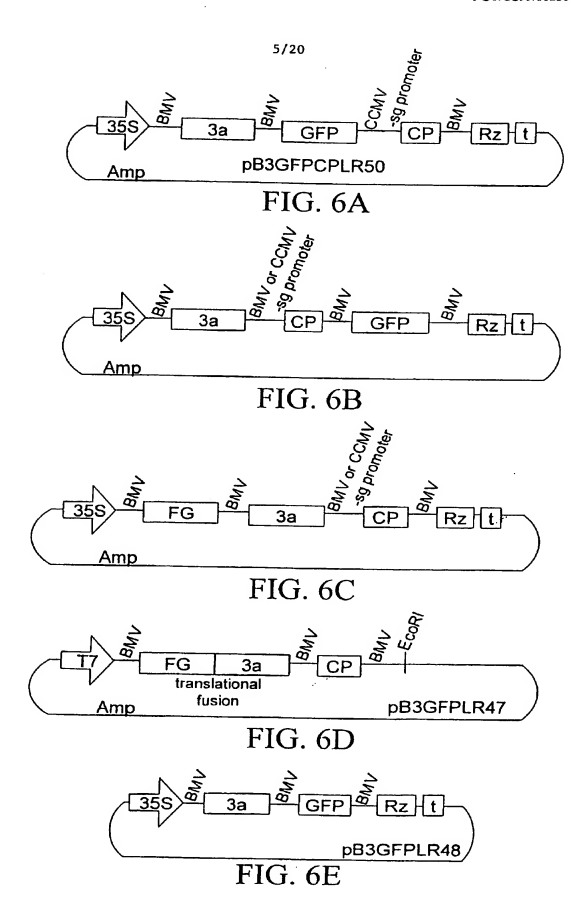
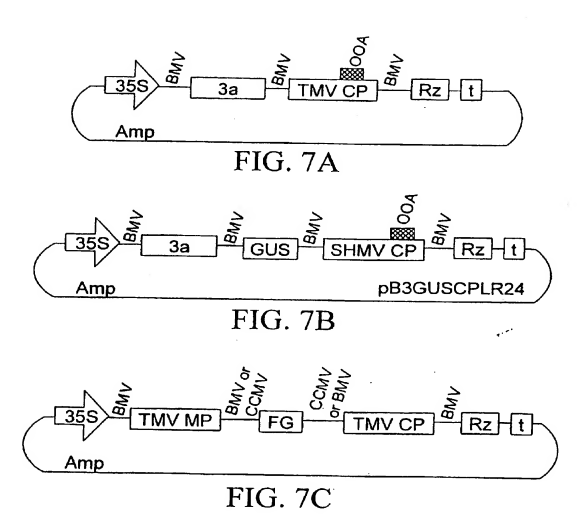


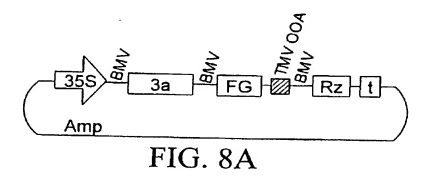
FIG. 5D

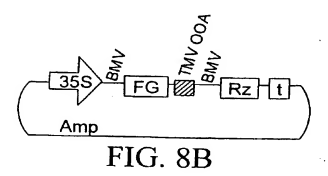


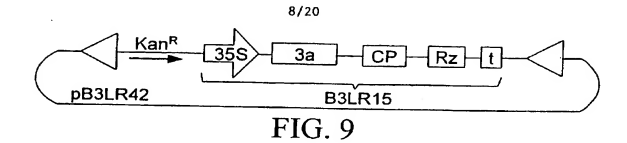
WO 99/61597

PCT/US99/11250









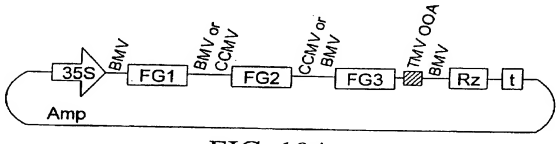
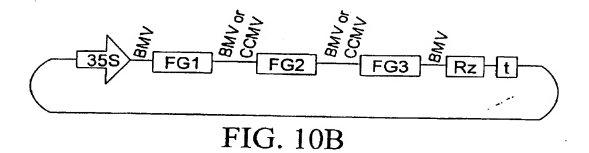
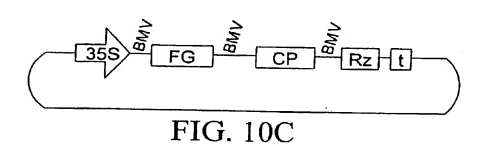
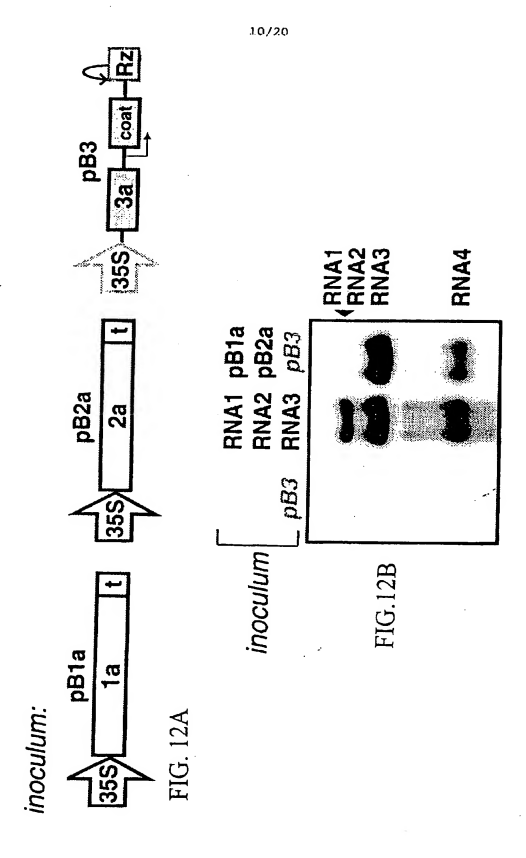


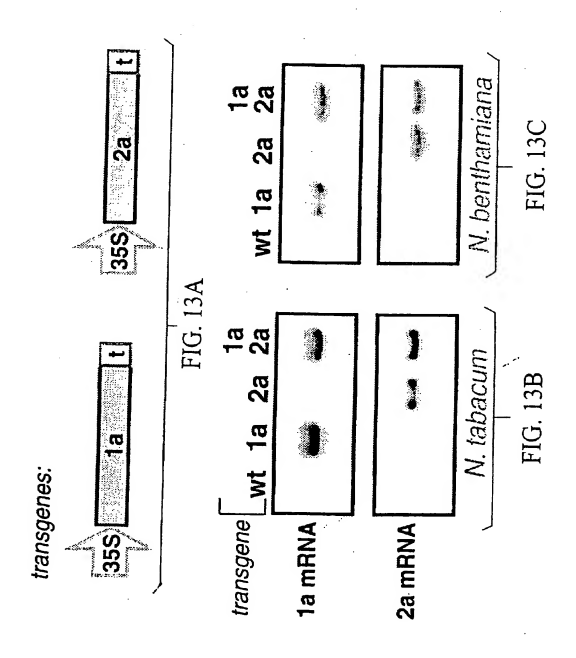
FIG. 10A



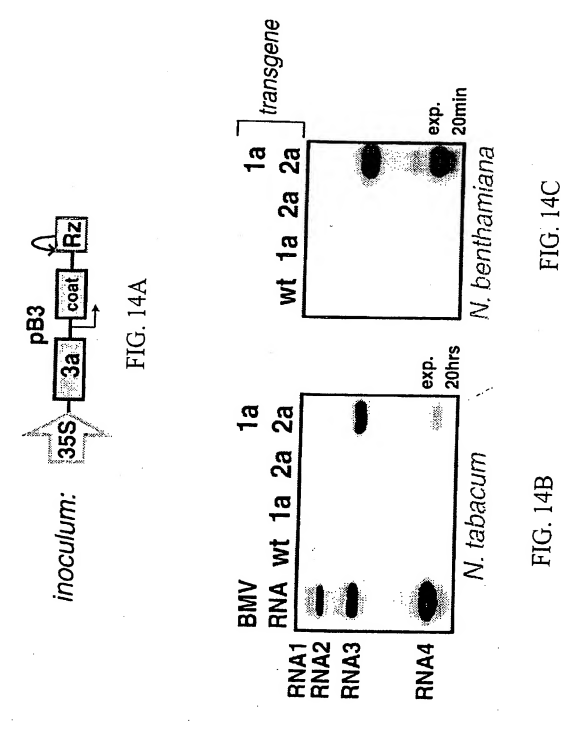


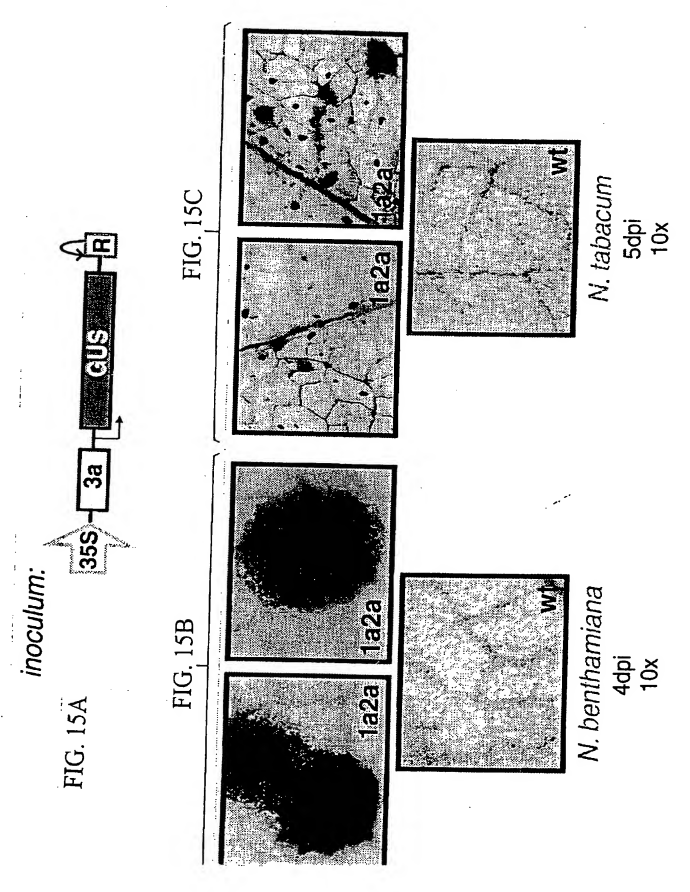






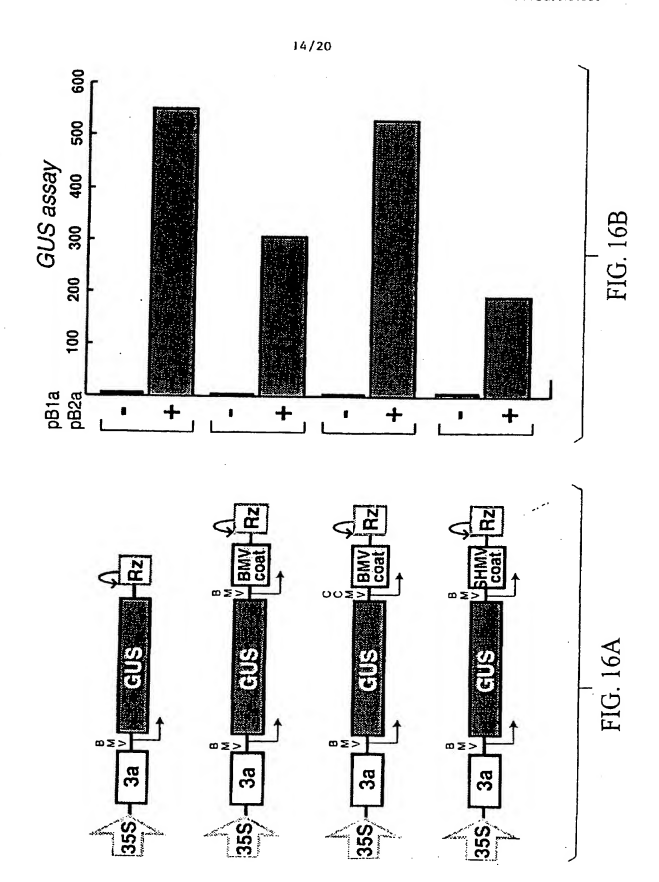
SUBSTITUTE SHEET (RULE 26)

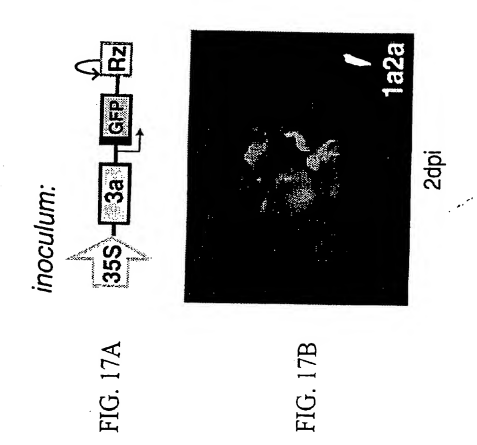


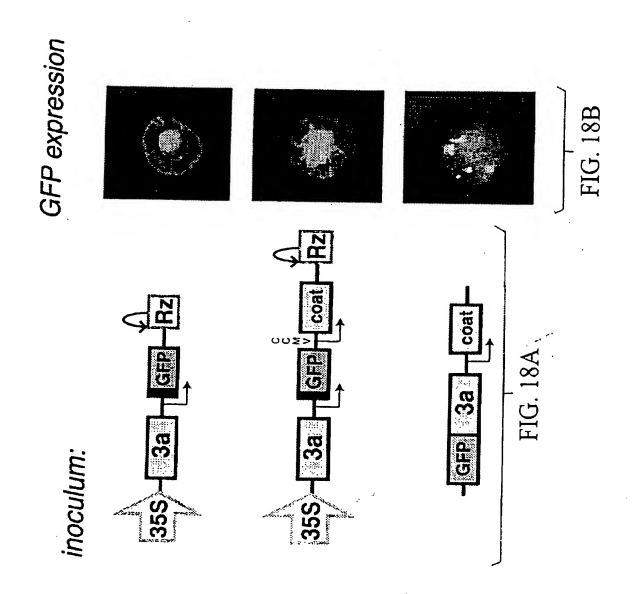


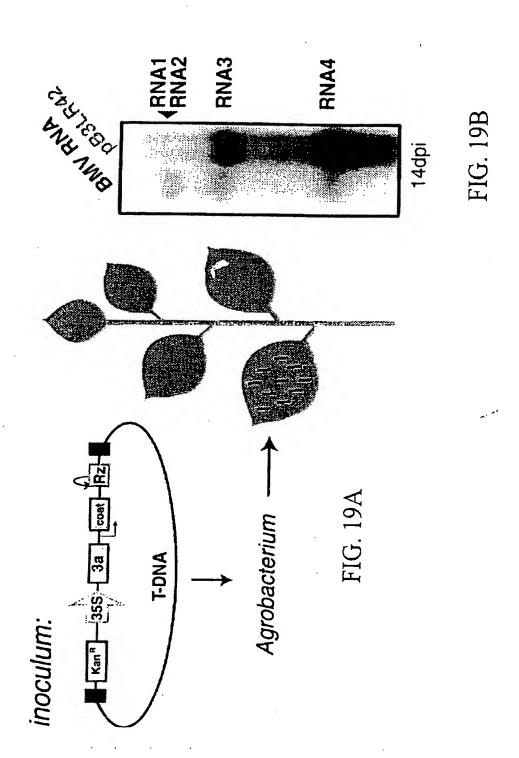
WO 99/61597

PCT/US99/11250









WO 99/61597

